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(54) Metal coordination compound, luminescene device and display apparatus

(57) An electroluminescence device having a layer containing a specific metal coordination compound is provided. The metal coordination compound is represented by formula (1) below:

$$ML_mL_n'$$
 (1),

wherein M is a metal atom of Ir, Pt, Rh or Pd; L and L' are mutually different bidentate ligands; m is 1, 2 or 3 and n is 0, 1 or 2 with the proviso that m+n is 2 or 3; a partial structure MLm is represented by formula (2) shown below and a partial structure ML'_n is represented by formula (3) or (4) shown below:

$$M = \begin{cases} CyN1 \\ CyC1 \end{cases} m \qquad (2) \qquad M = \begin{cases} CyN2 \\ CyC2 \end{cases} m \qquad (3) \qquad M = \begin{cases} CyN2 \\ O = S \\ O = \begin{cases} CyN2 \\ O = S \\ O$$

The metal coordination compound of the formula (1) is characterized by having at least one aromatic substituent for at least one of CyN1, CyN2, CyC1 and CyC2. The metal coordination compound having the aromatic substituent is effective in providing high-efficiency luminescence, long-term high luminance, and less deterioration by current passing.

Description

FIELD OF THE INVENTION AND RELATED ART

- [0001] The present invention relates to a luminescence device, a display apparatus and a metal coordination compound therefor. More specifically, the present invention relates to a luminescence device employing an organic metal coordination compound having a formula (1) appearing hereinafter as a luminescence material so as to allow stable luminescence efficiency, a display apparatus including the luminescence device and the metal coordination compound adapted for use in the luminescence device.
- [0002] An organic electroluminescence (EL) device has been extensively studied as a luminescence device with a high responsiveness and high efficiency.
 - [0003] The organic EL device generally has a sectional structure as shown in Figure 1A or 1B (e.g., as described in "Macromol. Symp.", 125, pp. 1 48 (1997)).
- [0004] Referring to the figures, the EL device generally has a structure including a transparent substrate 15, a transparent electrode 14 disposed on the transparent substrate 15, a metal electrode 11 disposed opposite to the transparent electrode 14, and a plurality of organic (compound) layers disposed between the transparent electrode 14 and the metal electrode 11.
 - [0005] Referring to Figure 1, the EL device in this embodiment has two organic layers including a luminescence layer 12 and a hole transport layer 13.
- [0006] The transparent electrode 14 may be formed of a film of ITO (indium tin oxide) having a larger work function to ensure a good hole injection performance into the hole transport layer. On the other hand, the metal electrode 11 may be formed of a layer of aluminum, magnesium, alloys thereof, etc., having a smaller work function to ensure a good electron injection performance into the organic layer(s).
 - [0007] These (transparent and metal) electrodes 14 and 11 may be formed in a thickness of 50 200 nm.
- 25 [0008] The luminescence layer 12 may be formed of, e.g., aluminum quinolinol complex (representative example thereof may include Alq3 described hereinafter) having an electron transporting characteristic and a luminescent characteristic. The hole transport layer 13 may be formed of, e.g., triphenyldiamine derivative (representative example thereof may include α-NPD described hereinafter) having an electron donating characteristic.
 - [0009] The above-described EL device exhibits a rectification characteristic, so that when an electric field is applied between the metal electrode 11 as a cathode and the transparent electrode 14 as an anode, electrons are injected from the metal electrode 11 into the luminescence layer 12 and holes are injected from the transparent electrodes 14.
 - [0010] The thus-injected holes and electrons are recombined within the luminescence layer 12 to produce excitons, thus causing luminescence. At that time, the hole transport layer 13 functions as an electron-blocking layer to increase a recombination efficiency at the boundary between the luminescence layer 12 and the hole transport layer 13, thus enhancing a luminescence efficiency.
 - [0011] Referring to Figure 1B, in addition to the layers shown in Figure 1A, an electron transport layer 16 is disposed between the metal electrode 11 and the luminescence layer 12, whereby an effective carrier blocking performance can be ensured by separating functions of luminescence, electron transport and hole transport, thus allowing effective luminescence.
- [0012] The electron transport layer 16 may be formed of, e.g., oxadiazole derivatives.

et al., Applied Physics Letters, Vol. 75, No. 1, pp. 4 - 6 (1999)).

- [0013] In ordinary organic EL devices, fluorescence caused during a transition of luminescent center molecule from a singlet excited state to a ground state is used as luminescence.
- [0014] On the other hand, not the above fluorescence (luminescence) via singlet exciton, phosphorescence (luminescence) via triplet exciton has been studied for use in organic EL device as described in, e.g., "Improved energy transfer in electrophosphorescent device" (D.F. O'Brien et al., Applied Physics Letters, Vol. 74, No. 3, pp. 442 444 (1999)) and "Very high-efficiency green organic light-emitting devices based on electrophosphorescence" (M.A. Baldo
- [0015] The EL devices shown in these documents may generally have a sectional structure shown in Figure 1C.
- [0016] Referring to Figure 1C, four organic layers including a hole transfer layer 13, a luminescence layer 12, an exciton diffusion-prevention layer 17, and an electron transport layer 16 are successively formed in this order on the transparent electrode (anode) 14.
 - [0017] In the above documents, higher efficiencies have been achieved by using four organic layers including a hole transport layer 13 of α-NPD (shown below), an electron transport layer 16 of Alq3 (shown below), an exciton diffusion-prevention layer 17 of BPC (shown below), and a luminescence layer 12 of a mixture of CPB (shown below) as a host material with Ir(ppy)₃ (shown below) or PtOEP (shown below) as a guest phosphorescence material doped into CBP at a concentration of ca. 6 wt. %.

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Alq3

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BCP

$$C_2H_5$$
 C_2H_5 C

Alq3: tris(8-hydroxyquinoline) aluminum (aluminum-quinolinol complex),

 α -NPD: N4,N4'-di-naphthalene-1-yl-N4,N4'-diphenyl-biphenyl-4,4'-diamine (4,4'-bis[N-(1-naphthyl)-N-phenyl-amino]biphenyl),

CBP: 4,4'-N,N'-dicarbazole-biphenyl,

BCP: 2,9-dimethyl-4,7-diphenyl-1,10-phenanthroline,

Ir(ppy)3: fac tris(2-phenylpyridine)iridium (iridium-phenylpyridine complex), and

PtEOP: 2,3,7,8,12,13,17,18-octaethyl-21H,23H-porphine platinum (platinum-octaethyl porphine complex).

[0018] The phosphorescence (luminescence) material used in the luminescence layer 12 has attracted notice. This

is because the phosphorescence material is expected to provide a higher luminescence efficiency in principle.

[0019] More specifically, in the case of the phosphorescence material, excitons produced by recombination of carriers comprise singlet excitons and triplet excitons presented in a ratio of 1:3. For this reason, when fluorescence caused during the transition from the singlet excited state to the ground state is utilized, a resultant luminescence efficiency is 25 % (as upper limit) based on all the produced excitons in principle.

[0020] On the other hand, in the case of utilizing phosphorescence caused during transition from the triplet excited state, a resultant luminescence efficiency is expected to be at least three times that of the case of fluorescence in principle. In addition thereto, if intersystem crossing from the singlet excited state (higher energy level) to the triplet excited state is taken into consideration, the luminescence efficiency of phosphorescence can be expected to be 100 % (four times that of fluorescence) in principle.

[0021] The use of phosphorescence based on transition from the triplet excited state has also been proposed in, e. g., Japanese Laid-Open Patent Application (JP-A) 11-329739, JP-A 11-256148 and JP-A 8-319482.

[0022] However, the above-mentioned organic EL devices utilizing phosphorescence have accompanied with a problem of luminescent deterioration particularly in an energized state.

[0023] The reason for luminescent deterioration has not been clarified as yet but may be attributable to such a phenomenon that the life of triplet exciton is generally longer than that of singlet exciton by at least three digits, so that molecule is placed in a higher-energy state for a long period to cause reaction with ambient substance, formation of exciplex or excimer, change in minute molecular structure, structural change of ambient substance, etc.

[0024] Accordingly, the (electro)phosphorescence EL device is expected to provide a higher luminescence efficiency as described above, while the EL device is required to suppress or minimize the luminescent deterioration in energized state. Further, a luminescence center material for the EL device is required to allow high-efficiency luminescence and exhibit a good stability.

SUMMARY OF THE INVENTION

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[0025] An object of the present invention is to provide a luminescence device capable of providing a high-efficiency luminescent state at a high brightness (or luminance) for a long period while minimizing the deterioration in luminescence in energized state.

[0026] Another object of the present invention is to provide a display apparatus including the luminescence device.

[0027] A further object of the present invention is to provide a metal coordination compound as a luminescence center material suitable for an organic layer for the luminescence device.

[0028] According to the present invention, there is provided a metal coordination compound (metal complex), particularly an iridium complex,

characterized by having at least one aromatic substituent. More specifically, there is provided a metal coordination compound represented by formula (1) below:

$$ML_mL_n'$$
 (1),

wherein M is a metal atom of Ir, Pt, Rh or Pd; L and L' are mutually different bidentate ligands; m is 1, 2 or 3 and n is 0, 1 or 2 with the proviso that m+n is 2 or 3; a partial structure MLm is represented by formula (2) shown below and a partial structure ML'_n is represented by formula (3) or (4) shown below:

wherein CyN1 and CyN2 are each cyclic group capable of having a substituent, including a nitrogen atom and bonded to the metal atom M via the nitrogen atom; CyC1 and CyC2 are each cyclic group capable of having a substituent, including a carbon atom and bonded to the metal atom M via the carbon atom with the proviso that the cyclic group CyN1 and the cyclic group CyN2 and the

cyclic group CyC2 are bonded to each other via a covalent bond;

the optional substituent of the cyclic groups is selected from a halogen atom, cyano group, a nitro group, a trialkylsilyl group of which the alkyl groups are independently a linear or branched alkyl group having 1 to 8 carbon atoms, a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -O-CO-, -CH=CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom; or an aromatic group capable of having a substituent which is selected from an aromatic group capable of having a substituent (that is a halogen atom, a cyano atom, a nitro atom, a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -CH=CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom), a halogen atom, a cyano atom, a nitro atom, and a linear or branched alkyl group having 1 to 20 carbon atoms (of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -CO-O-, -CH=CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom);

E and G are independently a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom, or an aromatic group capable of having a substituent (that is a halogen atom, a cyano atom, a nitro atom, a trialkylsilyl group of which the alkyl groups are independently a linear or branched alkyl group having 1 · 8 carbon atoms, a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -O-CO-, -CH=CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom; and

the cyclic groups CyN1, CyN2, CyC1 and CyC2 have at least one aromatic substituent capable of having a substituent which is selected from an aromatic group capable of having a substituent (that is a halogen atom, a cyano atom, a nitro atom, a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -O-CO-, -CH=CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom), a halogen atom, a cyano atom, a nitro atom, a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -O-CO-, -CH=CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom).

[0029] In the formula (1), M may preferably be Ir as described above, and n may preferably be 0.

[0030] In the formula (2), CyN1 and CyC1 may preferably be any one of the following combinations:

CyN1	CyC1
pyridyl pyridyl	naphthyl thienyl
pyridyl	benzothienyl

[0031] The present invention also provides an electroluminescence device, comprising: a pair of electrodes disposed on a substrate, and a luminescence unit comprising at least one organic compound disposed between the electrodes, wherein the organic compound comprises a metal coordination compound represented by the above-mentioned formula (1).

[0032] In the electroluminescence device, a voltage is applied between the electrodes to emit light.

[0033] In a preferred embodiment of the electroluminescence device, a voltage is applied between the electrodes to emit phosphorescence.

[0034] The present invention further provides a picture display apparatus, comprising an electroluminescence device described above and a means for supplying electric signals to the electroluminescence device.

[0035] These and other objects, features and advantages of the present invention will become more apparent upon a consideration of the following description of the preferred embodiments of the present invention taken in conjunction with the accompanying drawings.

BRIEF DESCRIPTION OF THE DRAWINGS

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Figures 1A, 1B and 1C illustrate embodiments of the luminescence device according to the present invention, respectively.

Figure 2 schematically illustrates a panel structure including an EL device and drive means.

Figures 3A, 3B and 3C show device performances of a luminescence device used in Example 9 appearing here-inafter, wherein Figure 3A shows an electric field strength-current density curve, Figure 3B shows an electric field strength-luminance curve, and Figure 3C shows a luminescence spectrum under application of a voltage of 10 volts.

DETAILED DESCRIPTION OF THE INVENTION

[0037] In the case where the luminescence layer comprises a host material having a carrier-transporting function and a phosphorescent guest material, a process of phosphorescence via triplet excitons may include unit processes as follows:

- 1. transportation of electrons and holes within a luminescence layer,
- 2. formation of host excitons,

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- 3. excitation energy transfer between host molecules,
- 4. excitation energy transfer from the host to the guest,
- 5. formation of guest triplet excitons, and
- 6. transition of the guest triplet excitons to the ground state and phosphorescence.

[0038] Desirable energy transfer in each unit process and luminescence are caused in competition with various energy deactivation processes.

[0039] Needless to say, a luminescence efficiency of an organic luminescence device is increased by increasing the luminescence quantum yield of a luminescence center material. In addition thereto, an efficient energy transfer between host material molecules and/or between host material molecule and guest material molecule is also an important factor.

[0040] Further, the above-described luminescent deterioration in energized state may presumably relate to the luminescent center material per se or an environmental change thereof by its ambient molecular structure.

[0041] For this reason, our research group has extensively investigated an effect of use of the metal coordination compound of formula (1) as the luminescent center material and as a result, has found that the metal coordination compound of formula (1) allows a high-efficiency luminescence with a high brightness (luminance) for a long period, and less deterioration in energized state.

[0042] The metal coordination compound represented by the above formula (1) according to the present invention causes phosphorescence (luminescence) and its lowest excited state is believed to be an MLT* (metal-to-ligand charge transfer) excited state or π - π * excited state in a triplet state. The phosphorescent emission of light (phosphorescence) is caused at the time of transition from such a state to the ground state.

[0043] The metal coordination compound of formula (1) according to the present invention has been found to provide a higher phosphorescence (quantum) yield of 0.05 - 0.9 and a shorter phosphorescence life of 1 - 40 μ sec, as a result of our luminescence experiment based on photoluminescence by photo-excitation.

[0044] The shorter phosphorescence life is necessary to provide a resultant EL device with a higher luminescence efficiency. This is because the longer phosphorescence life increases molecules placed in their triplet excited state which is a waiting state for phosphorescence, thus lowering the resultant luminescence efficiency particularly at a higher current density. Further, an emission wavelength can be controlled by changing appropriately substituents R1 to T6 and species of aromatic group of the metal coordination compound of the formula (1).

[0045] Also from these viewpoints, the metal coordination compound of formula (1) according to the present invention is a suitable luminescent material for an EL device with a higher phosphorescence yield and a shorter phosphorescence life.

[0046] Particularly, by providing an aromatic group as a substituent (i.e., aromatic substituent) of the metal coordination compound of the formula (1), the resultant substituent has π -electron system extended to the outside of the metal coordination compound molecules, thus facilitating energy transfer from a host material and assisting electron/hole transport functions to result in an improved carrier transport performance. Further, in the present invention, the metal coordination compound of the formula (1) may preferably have the cyclic group CyN1 and/or CyN2 having pyridine structure, a pyridine derivative wherein one of CH groups is substituted with N atom, and fine-membered ring structures containing nitrogen atom and/or sulfur atom. By these partial structures, the resultant metal coordination compound of the formula (1) can be synthesized with a high yield and an excellent stability necessary for the luminescence material. [0047] In addition, as substantiated in Examples appearing hereinafter, it has been confirmed that the metal coordination compound of the formula (1) also exhibited an excellent stability in a durability test by continuous current passage. This may be attributable to a controlled intermolecular interaction of the metal coordination compound of the present invention into the metal coordination compound thereby to change an intermolecular interaction. As a result, it becomes possible to suppress formation of exciton associates leading to thermal deactivation, thus also

reducing quenching process to improve phosphorescence yield and device characteristics.

[0048] In the present invention, as the aromatic substituent for the metal coordination compound of the formula (I), it is preferred to use an aromatic group selected from the group consisting of those (sPh to sPe) shown hereinafter.

[0049] In the present invention, the luminescence device may preferably include the organic layer comprising the above-mentioned metal coordination compound between a pair of oppositely disposed electrodes comprising a transparent electrode (anode) and a metal electrode (cathode) which are supplied with a voltage to cause luminescence, thus constituting an electric-field luminescence device.

[0050] The luminescence device of the present invention has a layer structure shown in Figures 1A to 1C as specifically described above.

[0051] By the use of the metal coordination compound of formula (1) of the present invention, the resultant luminescence device has a high luminescence efficiency as described above.

[0052] The luminescence device according to the present invention may be applicable to devices required to allow energy saving and high luminance, such as those for display apparatus and illumination apparatus, a light source for printers, and backlight (unit) for a liquid crystal display apparatus. Specifically, in the case of using the luminescence device of the present invention in the display apparatus, it is possible to provide a flat panel display apparatus capable of exhibiting an excellent energy saving performance, a high visibility and a good lightweight property. With respect to the light source, it becomes possible to replace a laser light source of laser beam printer currently used widely with the luminescence device according to the present invention. Further, when the luminescence device of the present invention is arranged in independently addressable arrays as an exposure means for effecting desired exposure of light to a photosensitive drum for forming an image, it becomes possible to considerably reducing the volume (size) of image forming apparatus. With respect to the illumination apparatus and backlight (unit), the resultant apparatus (unit) using the luminescence device of the present invention is expected to have an energy saving effect.

[0053] For the application to a display, a drive system using a thin-film transistor (TFT) drive circuit according to an active matrix-scheme may be used. Hereinbelow, an embodiment of using a device of the present invention in combination with an active matrix substrate is briefly described with reference to Figure 2.

[0054] Figure 2 illustrates an embodiment of panel structure comprising an EL device and drive means. The panel is provided with a scanning signal driver, a data signal driver and a current supply source which are connected to gate selection lines, data signal lines and current supply lines, respectively. At each intersection of the gate selection lines and the data signal lines, a display pixel electrode is disposed. The scanning signal drive sequentially selects the gate selection lines G1, G2, G3 ... Gn, and in synchronism herewith, picture signals are supplied from the data signal driver to display a picture (image).

[0055] By driving a display panel including a luminescence layer comprising a luminescence material of the present invention, it becomes possible to provide a display which exhibits a good picture quality and is stable even for a long period display.

[0056] Some synthetic paths for providing a metal coordination compound represented by the above-mentioned formula (1) are illustrated below with reference to an iridium coordination compound (m+n=3) for example:

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$$IrCl_3 \xrightarrow{2 \times L} [Ir(L)_2Cl]_2 \xrightarrow{L'} Ir(L)_2L'$$

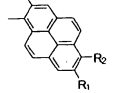
[0057] Other metal coordination compound (M = Pt, Rh and Pd) can also be synthesized in a similar manner.

[0058] Some specific structural examples of metal coordination compounds used in the present invention are shown in Tables 1 to Tables 17 appearing hereinafter, which are however only representative examples and are not exhaustive. Ph to sPe for CyN1, CyN2, CyC1, CyC2 and aromatic substituent(s) shown in Tables 1 to 17 represent partial structures shown below.

<CyC1, CyC2>

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$$R_1$$



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Cz:

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<CyN1, CyN2>

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<Aromatic substituent>

Table 1

B.A.		CVAN	CVC4	Ď	2	00	D.4
							R4
Ir	3	Pr	Ph	Н	Н	sPh	Н
lr	3	Pr	Ph	H	Η	sNp1	Ι
lr	з	Pr	Ph	Н	Η	sNp2	Н
lr	3	Pr	Ph	Н	Н	sTn1	Н
lr	3	Pr	Ph	Н	Н	sTn3	Н
lr	3	Pr	Ph	Н	Н	sPr	H
lr	3	Pr	Ph	Н	Н	sPe	Н
lr	3	Pr	Tn1	Н	Н	sPh	Н
lr	3	Pr	Tn1	Н	H	sNp1	Н
lr	3	Pr	Tn1	н	Н	sNp2	Н
lr	3	Pr	Tn1	Н	Н	sTn1	Н
lr	3	Pr	Tn1	Н	Н	sTn3	Н
Ir	3	Pr	⊤n1	н	Н	sPr	Н
lr	3	Pr	Tn1	Н	Н	sPe	Н
lr	3	Pr	Tn2	Н	Н	sPh	Н
lr	3	Pr	Tn2	Н	Н	sNp1	Н
lr	3	Pr	Tn2	Н	Н	sNp2	Н
lr	3	Pr	Tn2	Н	Н	sTn1	Н
lr	3	Pr	Tn2	Н	Н	sTn3	Н
lr	3	Pr	Tn2	Н	Н	sPr	Н
lr	3	Pr	Tn2	Н	Н	sPe	Н
lr	3	Pr	Tn3	Н	Н	sPh	Н
lr	3	Pr	Tn3	Н	Н	sNp1	Н
	Ir I	Ir 3 I	Ir 3 Pr Ir 3 Pr	Ir 3 Pr Ph Ir 3 Pr Tn1 Ir 3 Pr Tn2 Ir 3 Pr	Ir 3 Pr Ph H Ir 3 Pr Tn1 H Ir 3 Pr Tn2 H Ir 3 Pr Tn2 H Ir 3 Pr Tn2 H Ir 3 </td <td>Ir 3 Pr Ph H H Ir 3 Pr Tn1 H H Ir 3 Pr Tn2 H H <!--</td--><td>Ir 3 Pr Ph H H SPh Ir 3 Pr Ph H H sNp2 Ir 3 Pr Ph H H sNp2 Ir 3 Pr Ph H H sTn3 Ir 3 Pr Ph H H sPr Ir 3 Pr Ph H H sPr Ir 3 Pr Ph H H sPr Ir 3 Pr Tn1 H H sNp1 Ir 3 Pr Tn1 H H sNp1 Ir 3 Pr Tn1 H H sTn1 Ir 3 Pr Tn1 H H sPr Ir 3 Pr Tn1 H H sPr Ir 3 Pr Tn2 H H sNp1</td></td>	Ir 3 Pr Ph H H Ir 3 Pr Tn1 H H Ir 3 Pr Tn2 H H </td <td>Ir 3 Pr Ph H H SPh Ir 3 Pr Ph H H sNp2 Ir 3 Pr Ph H H sNp2 Ir 3 Pr Ph H H sTn3 Ir 3 Pr Ph H H sPr Ir 3 Pr Ph H H sPr Ir 3 Pr Ph H H sPr Ir 3 Pr Tn1 H H sNp1 Ir 3 Pr Tn1 H H sNp1 Ir 3 Pr Tn1 H H sTn1 Ir 3 Pr Tn1 H H sPr Ir 3 Pr Tn1 H H sPr Ir 3 Pr Tn2 H H sNp1</td>	Ir 3 Pr Ph H H SPh Ir 3 Pr Ph H H sNp2 Ir 3 Pr Ph H H sNp2 Ir 3 Pr Ph H H sTn3 Ir 3 Pr Ph H H sPr Ir 3 Pr Ph H H sPr Ir 3 Pr Ph H H sPr Ir 3 Pr Tn1 H H sNp1 Ir 3 Pr Tn1 H H sNp1 Ir 3 Pr Tn1 H H sTn1 Ir 3 Pr Tn1 H H sPr Ir 3 Pr Tn1 H H sPr Ir 3 Pr Tn2 H H sNp1

Table 1 (continued)

No	М	m	CyN1	CyC1	R1	R2	R3	R4
24	Ir	3	Pr	Tn3	Н	Н	sNp2	Н
25	łr	3	Pr	Tn3	Н	н	sTn1	Н
26	lr	3	Pr	Tn3	Н	н	sTn3	Н
27	lr	3	Pr	Tn3	н	Н	sPr	Н
28	lr	3	Pr	Tn3	н	Н	sPe	Н
29	lr	3	Pr	Tn4	Н	Н	sPh	Н
30	lr	3	Pr	Tn4	Н	Н	sNp1	Н
31	lr	3	Pr	Tn4	Н	Н	sNp2	Н
32	tr	3	Pr	Tn4	Н	Н	sTn1	Н
33	İr	3	Pr	Tn4	Н	н	sTn3	Н
34	lr	3	Pr	Tn4	Н	Н	sPr	Ι
35	lr	3	Pr	Tn4	Н	Н	sPe	H
36	lr	3	Pr	Np1	Н	Н	sPh	Н
37	lr	3	Pr	Np1	Н	н	sNp1	Н
38	lr	3	Pr	Np1	Н	Н	sNp2	Н
39	lr	3	Pr	Np1	Н	Н	sTn1	Н
40	Ir	3	Pr	Np1	Н	Н	sTn3	Н
41	lr	3	Pr	Np1	Н	Н	sPr	Н
42	lr	3	Pr	Np1	Н	Н	sPe	Н
43	lr	3	Pr	Np2	Н	H	Н	sPh
44	Ir	3	Pr	Np2	Н	Н	sNp1	Н
45	Ir	3	Pr	Np2	Н	Н	sNp2	Н
46	lr	ფ	Pr	Np2	Н	Н	sTn1	Н
47	lr	з	Pr	Np2	Н	Н	sTn3	Н
48	lr	3	Pr	Np2	Н	Н	sPr	Η
49	lr	3	Pr	Np2	Н	Н	sPe	Η
50	lr	3	Pr	Pe	Н	Н	sPh	H
51	lr	3	Pr	Pe	Н	н	sNp1	Н
52	lr	3	Pr	Pe	Н	Н	sNp2	I

Table 2

No	Σ	E	CyN1	CyC1	R1	R2	R3	R4
53	lr	3	Pr	Pe	Н	н	sTn1	н
54	lr	3	Pr	Pe	Н	Н	sTn3	Н
55	lr	3	Pr	Pe	Н	н	sPr	Н
56	lr	3	Pr	Pe	Н	Н	sPe	Н
57	lr	3	Pr	Cn1	Н	Н	sPh	Н
58	lr	3	Pr	Cn1	Н	Н	sNp1	Н

Table 2 (continued)

				14510	Z (COIIII	1404			
	No	М	m	CyN1	CyC1	R1	R2	R3	R4
	59	lr	3	Pr	Cn1	Н	Η	sNp2	Н
	60	Ir	3	Pr	Cn1	Н	Н	sTn1	Н
	61	lr	3	Pr	Cn1	H	Н	sTn3	Н
	62	lr	3	Pr	Cn1	Н.	H	sPr	Н
	63	lr	3	Pr	Cn1	Н	Η	sPe	Н
	64	lr	3	Pr	Cn2	Н	Ι	sPh	Н
	65	lr	3	Pr	Cn2	Н	I	sNp1	Н
	66	lr	3	. Pr	Cn2	Н	Η	sNp2	н
•	67	lr	3	Pr	Cn2	Н	Ι	sTn1	Н
•	68	lr	3	Pr	Cn2	Н	Н	sTn3	Н
•	69	Ir	3	Pr	Cn2	Н	I	sPr	Н
•	70	lr	3	Pr	Cn2	Н	Η	sPe	Н
	71	Ir	3	Pr	Cz	Н	Н	sPh	Η
•	72	lr	3	Pr	Cz	Н	Ι	sNp1	Н
	73	lr	3	Pr	Cz	н	Η	sNp2	Н
	74	lr	3	Pr	Cz	Н	Η	sTn1	Н
	75	lr	3	Pr	Cz	Н	Τ	sTn3	Н
	76	Ir	3	Pr	Cz	Н	Ħ	sPr	Н
	77	Ir	3	Pr	Cz	Н	Н	sPe	Н
	78	lr	3	Pd	Ph	Н	Н	sPh	Н
	79	lr	3	Pd	Ph	Н	Н	sNp1	Н
	80	Ir	3	Pd	Ph	Н	Η	sNp2	Н
	81	Ir	3	Pd	Ph	Н	Н	sTn1	Н
	82	Ir	3	Pd	Ph	Н	Н	sTn3	Н
	83	lr	3	Pd	Ph	н	Н	sPr	Н
	84	lr	3	Pd	Ph	н	Н	sPe	Н
	85	lr	3	Pd	Tn1	Н	Н	sPh	Н
	86	lr.	3	Pd	Tn1	Н	Н	sNp1	Н
	87	lr	3	Pd	Tn1	Н	Н	sNp2	Н
	88	lr	3	Pd	Tn1	Н	Н	sTn1	Н
	89	lr	3	Pd	Tn1	Н	Н	sTn3	Н
	90	lr	3	Pd	Tn1	Н	Н	sPr	Н
	91	lr	3	Pd	Tn1	Н	Н	sPe	н
	92	lr	3	Pd	Tn2	Н	Н	sPh	Н
	93	lr	3	Pd	Tn2	Н	Н	sNp1	Н
	94	Ir	3	Pd	Tn2	Н	Н	sNp2	н
	95	Ir	3	Pd	Tn2	Н	Н	sTn1	Н
	96	tr	3	Pd	Tn2	Н	Н	sTn3	Н
					·				

Table 2 (continued)

No	Δ	E	CyN1	CyC1	R1	R2	R3	R4
97	lr	3	Pd	Tn2	Н	Н	sPr	Н
98	lr	3	Pd	Tn2	Н	Н	sPe	н
99	lr	3	Pd	Tn3	Н	Н	sPh	Н
100	lr	3	Pd	Tn3	Н	Н	sNp1	Н
101	lr	3	Pd	Tn3	Н	Н	sNp2	Н
102	lr	3	Pd	Tn3	Н	Н	sTn1	Н
103	lr	3	Pd	Tn3	Н	Н	sTn3	H
104	lr	3	Pd	Tn3	Н	Н	sPr	Н

Table 3

No M m CyN1 CyC1 R1 R2 R3 R4 105 Ir 3 Pd Tn3 H H sPe H 106 Ir 3 Pd Tn4 H H sPh H 107 Ir 3 Pd Tn4 H H sNp1 H 108 Ir 3 Pd Tn4 H H sNp2 H 109 Ir 3 Pd Tn4 H H sTn1 H 109 Ir 3 Pd Tn4 H H sTn3 H 110 Ir 3 Pd Tn4 H H sTn3 H 111 Ir 3 Pd Tn4 H H sPr H 112 Ir 3 Pd Np1 H H sNp1 H 113 Ir	Table 3										
106 Ir 3 Pd Tn4 H H sPh H 107 Ir 3 Pd Tn4 H H sNp1 H 108 Ir 3 Pd Tn4 H H sNp2 H 109 Ir 3 Pd Tn4 H H sTn1 H 110 Ir 3 Pd Tn4 H H sTn3 H 111 Ir 3 Pd Tn4 H H sPr H 112 Ir 3 Pd Tn4 H H sPr H 113 Ir 3 Pd Np1 H H sPr H 114 Ir 3 Pd Np1 H H sNp1 H 115 Ir 3 Pd Np1 H H sNp2 H 116 Ir	No	М	m	CyN1	CyC1	R1	R2	R3	R4		
107 Ir 3 Pd Tn4 H H sNp1 H 108 Ir 3 Pd Tn4 H H sNp2 H 109 Ir 3 Pd Tn4 H H sTn1 H 110 Ir 3 Pd Tn4 H H sTn3 H 111 Ir 3 Pd Tn4 H H sSPr H 111 Ir 3 Pd Tn4 H H sSPr H 111 Ir 3 Pd Np1 H H sSPr H 112 Ir 3 Pd Np1 H H sNp1 H 114 Ir 3 Pd Np1 H H sNp2 H 115 Ir 3 Pd Np1 H H sSPr H 116 Ir	105	lr	3	Pd	Tn3	Н	Н	sPe	Η		
108 Ir 3 Pd Tn4 H H sNp2 H 109 Ir 3 Pd Tn4 H H sTn1 H 110 Ir 3 Pd Tn4 H H sTn3 H 111 Ir 3 Pd Tn4 H H sPr H 112 Ir 3 Pd Tn4 H H sPr H 112 Ir 3 Pd Np1 H H sPp H 113 Ir 3 Pd Np1 H H sNp1 H 114 Ir 3 Pd Np1 H H sNp2 H 115 Ir 3 Pd Np1 H H sNp2 H 116 Ir 3 Pd Np1 H H sTn3 H 117 Ir	106	lr	3	Pd	Tn4	Н	Η	sPh	Η		
109 Ir 3 Pd Tn4 H H sTn1 H 110 Ir 3 Pd Tn4 H H sTn3 H 111 Ir 3 Pd Tn4 H H sPr H 112 Ir 3 Pd Tn4 H H sPr H 112 Ir 3 Pd Np1 H H sPe H 113 Ir 3 Pd Np1 H H sPp H 114 Ir 3 Pd Np1 H H sNp2 H 115 Ir 3 Pd Np1 H H sNp2 H 116 Ir 3 Pd Np1 H H sNp2 H 115 Ir 3 Pd Np1 H H sTn3 H 119 Ir	107	lr	3	Pd	Tn4	Н	Н	sNp1	Η		
110 Ir 3 Pd Tn4 H H sTn3 H 111 Ir 3 Pd Tn4 H H sPr H 112 Ir 3 Pd Tn4 H H sPr H 113 Ir 3 Pd Np1 H H sPp H 114 Ir 3 Pd Np1 H H sNp1 H 115 Ir 3 Pd Np1 H H sNp2 H 116 Ir 3 Pd Np1 H H sNp2 H 116 Ir 3 Pd Np1 H H sTn3 H 116 Ir 3 Pd Np1 H H sTn3 H 117 Ir 3 Pd Np1 H H sPn H 119 Ir	108	lr	3	Pd	Tn4	Н	Н	sNp2	I		
1111 Ir 3 Pd Tn4 H H sPr H 112 Ir 3 Pd Tn4 H H sPe H 113 Ir 3 Pd Np1 H H sPh H 114 Ir 3 Pd Np1 H H sNp1 H 115 Ir 3 Pd Np1 H H sNp2 H 116 Ir 3 Pd Np1 H H sTn1 H 117 Ir 3 Pd Np1 H H sTn3 H 118 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np2 H H sNp1 H 120 Ir	109	lr	3	Pd	Tn4	Н	Н	sTn1	Н		
112 Ir 3 Pd Tn4 H H sPe H 113 Ir 3 Pd Np1 H H sPh H 114 Ir 3 Pd Np1 H H sNp1 H 115 Ir 3 Pd Np1 H H sNp2 H 116 Ir 3 Pd Np1 H H sNp2 H 116 Ir 3 Pd Np1 H H sTn1 H 116 Ir 3 Pd Np1 H H sTn1 H 117 3 Pd Np1 H H sTn3 H 118 Ir 3 Pd Np1 H H sPe H 119 Ir 3 Pd Np2 H H sPp H 120 Ir 3 <	110	lr	3	Pd	Tn4	Н	Н	sTn3	Η		
113 Ir 3 Pd Np1 H H sPh H 114 Ir 3 Pd Np1 H H sNp1 H 115 Ir 3 Pd Np1 H H sNp2 H 116 Ir 3 Pd Np1 H H sNp2 H 117 Ir 3 Pd Np1 H H sTn3 H 118 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np2 H H sPr H 120 Ir 3 Pd Np2 H H sNp1 H 123 Ir <	111	lr	3	Pd	Tn4	Н	Н	sPr	H		
114 Ir 3 Pd Np1 H H sNp1 H 115 Ir 3 Pd Np1 H H sNp2 H 116 Ir 3 Pd Np1 H H sTn1 H 117 Ir 3 Pd Np1 H H sTn3 H 118 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np2 H H sPr H 119 Ir 3 Pd Np2 H H sPr H 119 Ir 3 Pd Np2 H H sNp1 H 120 Ir 3 Pd Np2 H H sTn1 H 123 Ir	112	lr	3	Pd	Tn4	Н	Н	sPe	н		
115 Ir 3 Pd Np1 H H sNp2 H 116 Ir 3 Pd Np1 H H sTn1 H 117 Ir 3 Pd Np1 H H sTn3 H 118 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np2 H H sPh H 120 Ir 3 Pd Np2 H H sPh H 121 Ir 3 Pd Np2 H H sNp1 H 122 Ir 3 Pd Np2 H H sTn1 H 123 Ir 3 Pd Np2 H H sTn3 H 125 Ir	113	lr	3	Pd	Np1	Н	Н	sPh	Н		
116 Ir 3 Pd Np1 H H sTn1 H 117 Ir 3 Pd Np1 H H sTn3 H 118 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np2 H H sPr H 120 Ir 3 Pd Np2 H H sPn H 121 Ir 3 Pd Np2 H H sNp1 H 122 Ir 3 Pd Np2 H H sNp2 H 123 Ir 3 Pd Np2 H H sTn3 H 124 Ir 3 Pd Np2 H H sPr H 125 Ir <	114	lr	3	Pd	Np1	Н	Н	sNp1	H		
117 Ir 3 Pd Np1 H H sTn3 H 118 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np1 H H sPr H 120 Ir 3 Pd Np2 H H sPh H 121 Ir 3 Pd Np2 H H sNp1 H 122 Ir 3 Pd Np2 H H sNp2 H 123 Ir 3 Pd Np2 H H sTn1 H 124 Ir 3 Pd Np2 H H sTn3 H 125 Ir 3 Pd Np2 H H sPr H 126 Ir 3 Pd Pe H H sPp H 127 Ir <t< td=""><td>115</td><td>lr</td><td>3</td><td>Pd</td><td>Np1</td><td>Н</td><td>Н</td><td>sNp2</td><td>H</td></t<>	115	lr	3	Pd	Np1	Н	Н	sNp2	H		
118 Ir 3 Pd Np1 H H sPr H 119 Ir 3 Pd Np1 H H sPe H 120 Ir 3 Pd Np2 H H sPh H 121 Ir 3 Pd Np2 H H sNp1 H 122 Ir 3 Pd Np2 H H sNp2 H 123 Ir 3 Pd Np2 H H sTn1 H 124 Ir 3 Pd Np2 H H sTn3 H 125 Ir 3 Pd Np2 H H sPr H 126 Ir 3 Pd Np2 H H sPe H 127 Ir 3 Pd Pe H H sNp1 H 128 Ir <t< td=""><td>116</td><td>lr</td><td>3</td><td>Pd</td><td>Np1</td><td>Н</td><td>Н</td><td>sTn1</td><td>Н</td></t<>	116	lr	3	Pd	Np1	Н	Н	sTn1	Н		
119 Ir 3 Pd Np1 H H sPe H 120 Ir 3 Pd Np2 H H sPh H 121 Ir 3 Pd Np2 H H sNp1 H 122 Ir 3 Pd Np2 H H sNp2 H 123 Ir 3 Pd Np2 H H sTn1 H 124 Ir 3 Pd Np2 H H sTn3 H 125 Ir 3 Pd Np2 H H sPr H 126 Ir 3 Pd Np2 H H sPr H 127 Ir 3 Pd Pe H H sNp1 H 128 Ir 3 Pd Pe H H sNp1 H 129 Ir <t< td=""><td>117</td><td>lr</td><td>3</td><td>Pdi</td><td>Np1</td><td>Н</td><td>Н</td><td>sTn3</td><td>н</td></t<>	117	lr	3	Pdi	Np1	Н	Н	sTn3	н		
120 Ir 3 Pd Np2 H H sPh H 121 Ir 3 Pd Np2 H H sNp1 H 122 Ir 3 Pd Np2 H H sNp2 H 123 Ir 3 Pd Np2 H H sTn1 H 124 Ir 3 Pd Np2 H H sTn3 H 125 Ir 3 Pd Np2 H H sPr H 126 Ir 3 Pd Np2 H H sPr H 127 Ir 3 Pd Pe H H sNp1 H 128 Ir 3 Pd Pe H H sNp1 H 129 Ir 3 Pd Pe H H sNp2 H 130 Ir <t< td=""><td>118</td><td>lr</td><td>3</td><td>Pd</td><td>Np1</td><td>Н</td><td>Н</td><td>sPr</td><td>H</td></t<>	118	lr	3	Pd	Np1	Н	Н	sPr	H		
121 Ir 3 Pd Np2 H H sNp1 H 122 Ir 3 Pd Np2 H H sNp2 H 123 Ir 3 Pd Np2 H H sTn1 H 124 Ir 3 Pd Np2 H H sTn3 H 125 Ir 3 Pd Np2 H H sPr H 126 Ir 3 Pd Np2 H H sPe H 127 Ir 3 Pd Pe H H sNp1 H 128 Ir 3 Pd Pe H H sNp1 H 129 Ir 3 Pd Pe H H sNp2 H 130 Ir 3 Pd Pe H H sTn1 H	119	lr	3	Pd	Np1	Н	Н	sPe	Н		
122 Ir 3 Pd Np2 H H sNp2 H 123 Ir 3 Pd Np2 H H sTn1 H 124 Ir 3 Pd Np2 H H sTn3 H 125 Ir 3 Pd Np2 H H sPr H 126 Ir 3 Pd Np2 H H sPe H 127 Ir 3 Pd Pe H H sPh H 128 Ir 3 Pd Pe H H sNp1 H 129 Ir 3 Pd Pe H H sNp2 H 130 Ir 3 Pd Pe H H sTn1 H	120	lr	3	Pd	Np2	Н	Н	sPh	Н		
123 Ir 3 Pd Np2 H H sTn1 H 124 Ir 3 Pd Np2 H H sTn3 H 125 Ir 3 Pd Np2 H H sPr H 126 Ir 3 Pd Np2 H H sPe H 127 Ir 3 Pd Pe H H sPh H 128 Ir 3 Pd Pe H H sNp1 H 129 Ir 3 Pd Pe H H sNp2 H 130 Ir 3 Pd Pe H H sTn1 H	121	lr	3	Pd	Np2	Н	Н	sNp1	Н		
124 Ir 3 Pd Np2 H H sTn3 H 125 Ir 3 Pd Np2 H H sPr H 126 Ir 3 Pd Np2 H H sPe H 127 Ir 3 Pd Pe H H sPh H 128 Ir 3 Pd Pe H H sNp1 H 129 Ir 3 Pd Pe H H sNp2 H 130 Ir 3 Pd Pe H H sTn1 H	122	lr	3	Pd	Np2	Н	Н	sNp2	Н		
125 Ir 3 Pd Np2 H H sPr H 126 Ir 3 Pd Np2 H H sPe H 127 Ir 3 Pd Pe H H sPh H 128 Ir 3 Pd Pe H H sNp1 H 129 Ir 3 Pd Pe H H sNp2 H 130 Ir 3 Pd Pe H H sTn1 H	123	lr	3	Pd	Np2	Н	H	sTn1	Н		
126 Ir 3 Pd Np2 H H sPe H 127 Ir 3 Pd Pe H H sPh H 128 Ir 3 Pd Pe H H sNp1 H 129 Ir 3 Pd Pe H H sNp2 H 130 Ir 3 Pd Pe H H sTn1 H	124	lr	3	Pd	Np2	Н	Н	sTn3	Н		
127 Ir 3 Pd Pe H H sPh H 128 Ir 3 Pd Pe H H sNp1 H 129 Ir 3 Pd Pe H H sNp2 H 130 Ir 3 Pd Pe H H sTn1 H	125	lr	3	Pd	Np2	Н	н	sPr	Н		
128 Ir 3 Pd Pe H H sNp1 H 129 Ir 3 Pd Pe H H sNp2 H 130 Ir 3 Pd Pe H H sTn1 H	126	lr	3	Pd	Np2	Н	Н	sPe	Н		
129 Ir 3 Pd Pe H H sNp2 H 130 Ir 3 Pd Pe H H sTn1 H	127	lr	3	Pd	Pe	Н	Н	sPh	Н		
130 Ir 3 Pd Pe H H sTn1 H	128	lr	3	Pd	Pe	Н	Н	sNp1	Н		
	129	lr	3	Pd	Pe	Н	Н	sNp2	Н		
	130	lr	3	Pd	Pe	Н	Н	sTn1	Н		
131 Ir 3 Pd Pe H H sTn3 H	131	lr	3	Pd	Pe	Н	Н	sTn3	Н		

Table 3 (continued)

No	М	m	CyN1	CyC1	R1	R2	R3	R4
132	lr	3	Pd	Pe	Н	Н	sPr	Ι
133	lr	3	Pd	Pe	Н	Н	sPe	Н
134	lr	3	Pd	Cn1	Н	Н	sPh	Н
135	lr	3	Pd	Cn1	Н	н	sNp1	Н
136	lr	3	Pd	Cn1	н	Н	sNp2	Н
137	lr	3	Pd	Cn1	Н	Н	sTn1	H
138	ir	3	Pd	Cn1	Н	Н	sTn3	Н
139	lr	3	Pd	Cn1	Н	Н	sPr	Н
140	lr	3	Pd	Cn1	Н	Н	sPe	Н
141	lr	3	Pd	Cn2	Н	Н	sPh	Н
142	lr	3	Pd	Cn2	Н	Н	sNp1	Н
143	ir	3	Pd	Cn2	Н	Н	sNp2	Н
144	lr	3	Pd	Cn2	Н	Н	sTn1	Н
145	lr	3	Pd	Cn2	Н	Н	sTn3	Н
146	lr	3	Pd	Cn2	Н	Ι	sPr	H
147	lr	3	Pd	Cn2	Н	Н	sPe	Н
148	lr	3	Pd	Cz	Н	Н	sPh	H
149	lr	3	Pd	Cz	Н	Н	sNp1	Н
150	lr	3	Pd	Cz	Н	Н	sNp2	Н
151	lr	3	Pd	Cz	Н	Н	sTn1	Н
152	lr	3	Pd	Cz	Н	Н	sTn3	Н
153	lr	3	Pd	Cz	Н	Н	sPr	Н
154	Ir	3	Pd	Cz	Н	Н	sPe	H
155	lr	3	Pz	Ph	Н	Н	sPh	Н
156	lr	3	Pd	Ph	Н	Н	sNp1	Н

Table 4

No	М	m	CyN1	CyC1	R1	R2	R3	R4
157	lr	3	Pd	Ph	Н	Н	sNp2	Н
158	Ir	3	Pd	Ph	Н	Н	sTn1	Н
159	lr	3	Pd	Ph	Н	Н	sTn3	Η
160	lr	3	Pd	Ph	Н	Н	sPr	Н
161	lr	3	Pd	Ph	Н	Н	sPe	Н
162	lr	3	Pd	Tn1	Н	Ι	sPh	Н
163	lr	3	Pd	Tn1	H	Ι	sNp1	Ι
164	lr	3	Pd	Tn1	Н	Н	sNp2	H
165	lr	3	Pd	Tn1	Н	Н	sTn1	Н
166	lr	3	Pd	Tn1	Н	Н	sTn3	Н

Table 4 (continued)

No M m CyN1 CyC1 R1 R2 R3 R4 167 Ir 3 Pd Tn1 H H sPr H 168 Ir 3 Pd Tn1 H H sPr H 169 Ir 3 Pd Tn2 H H sNp1 H 170 Ir 3 Pd Tn2 H H sNp1 H 171 Ir 3 Pd Tn2 H H sNp2 H 172 Ir 3 Pd Tn2 H H sTn3 H 173 Ir 3 Pd Tn2 H H sFn3 H 174 Ir 3 Pd Tn3 H H sFn3 H 175 Ir 3 Pd Tn3 H H sNp1 H 18 Ir 3										
168 Ir 3 Pd Tn1 H H sPe H 169 Ir 3 Pd Tn2 H H sPh H 170 Ir 3 Pd Tn2 H H sNp1 H 171 Ir 3 Pd Tn2 H H sNp2 H 172 Ir 3 Pd Tn2 H H sTn1 H 173 Ir 3 Pd Tn2 H H sTn1 H 174 Ir 3 Pd Tn2 H H sPp H 175 Ir 3 Pd Tn2 H H sPp H 176 Ir 3 Pd Tn3 H H sPp H 177 Ir 3 Pd Tn3 H H sNp1 H 178 Ir <		No	М	m	CyN1	CyC1	R1	R2	R3	R4
169 Ir 3 Pd Tn2 H H sPh H 170 Ir 3 Pd Tn2 H H sNp1 H 171 Ir 3 Pd Tn2 H H sNp2 H 172 Ir 3 Pd Tn2 H H sNp2 H 173 Ir 3 Pd Tn2 H H sSPn H 174 Ir 3 Pd Tn2 H H sPp H 175 Ir 3 Pd Tn3 H H sPp H 176 Ir 3 Pd Tn3 H H sNp1 H 177 Ir 3 Pd Tn3 H H sNp2 H 178 Ir 3 Pd Tn3 H H sPn H H H sPn <		167	!r	3	Pd	Tn1	Н	Н	sPr	Н
170 Ir 3 Pd Tn2 H H SNp1 H 171 Ir 3 Pd Tn2 H H SNp2 H 172 Ir 3 Pd Tn2 H H SNp2 H 172 Ir 3 Pd Tn2 H H STn1 H 173 Ir 3 Pd Tn2 H H SFn H 175 Ir 3 Pd Tn2 H H SPr H 175 Ir 3 Pd Tn2 H H SPr H 176 Ir 3 Pd Tn3 H H SPp H 177 Ir 3 Pd Tn3 H H SNp1 H 178 Ir 3 Pd Tn3 H H SNp1 H 178 Ir 3 Pd Tn3 H H SNp2 H 179 Ir 3 Pd Tn3 H H STn3 H 180 Ir 3 Pd Tn3 H H STn3 H 181 Ir 3 Pd Tn3 H H SPr H 182 Ir 3 Pd Tn3 H H SPr H 182 Ir 3 Pd Tn3 H H SPr H 182 Ir 3 Pd Tn3 H H SPr H 184 Ir 3 Pd Tn3 H H SPr H 185 Ir 3 Pd Tn4 H H SNp1 H 185 Ir 3 Pd Tn4 H H SNp1 H 186 Ir 3 Pd Tn4 H H SNp1 H 187 Ir 3 Pd Tn4 H H STn3 H 188 Ir 3 Pd Tn4 H H STn3 H 188 Ir 3 Pd Tn4 H H SPr H 189 Ir 3 Pd Tn4 H H SPr H 189 Ir 3 Pd Tn4 H H SPr H 190 Ir 3 Pd Tn4 H H SPr H 191 Ir 3 Pd Tn4 H H SPr H 192 Ir 3 Pd Np1 H H SPr H 194 Ir 3 Pd Np1 H H SPr H 195 Ir 3 Pd Np1 H H SPr H 196 Ir 3 Pd Np1 H H SPr H 196 Ir 3 Pd Np1 H H SPr H 196 Ir 3 Pd Np1 H H SPr H 196 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199 Ir 3 Pd Np2 H H SPr H 199		168	lr	3	Pd	Tn1	Н	Н	sPe	н
171		169	lr	3	Pd	Tn2	Н	Н	sPh	Н
172		170	lr	3	Pd	Tn2	Н	Н	sNp1	Н
173 Ir 3 Pd Tn2 H H spn spn 172 H H spn H 175 H H spn H		171	Ir	3	Pd	Tn2	Н	Н	sNp2	Н
174 Ir 3 Pd Tn2 H H sPr H 175 Ir 3 Pd Tn2 H H sPr H 176 Ir 3 Pd Tn3 H H sPr H 177 Ir 3 Pd Tn3 H H sNp1 H 178 Ir 3 Pd Tn3 H H sNp2 H 179 Ir 3 Pd Tn3 H H sNp2 H 180 Ir 3 Pd Tn3 H H sSp7 H 181 Ir 3 Pd Tn3 H H sSp7 H 182 Ir 3 Pd Tn3 H H sSp7 H 182 Ir 3 Pd Tn4 H sSp7 H 182 Ir 3		172	lr	3	Pd	Tn2	I	Н	sTn1	Н
175 Ir 3 Pd Tn2 H H sPe H 176 Ir 3 Pd Tn3 H H sPh H 177 Ir 3 Pd Tn3 H H sNp1 H 178 Ir 3 Pd Tn3 H H sNp2 H 179 Ir 3 Pd Tn3 H H sNp2 H 180 Ir 3 Pd Tn3 H H sSFn3 H 181 Ir 3 Pd Tn3 H H sPr H 182 Ir 3 Pd Tn3 H H sPr H 182 Ir 3 Pd Tn4 H sPr H 182 Ir 3 Pd Tn4 H sPr H 183 Ir 3 Pd		173	lr	3	Pď	Tn2	Н	Н	sTn3	Н
176 Ir 3 Pd Tn3 H H sNp1 H 177 Ir 3 Pd Tn3 H H sNp1 H 178 Ir 3 Pd Tn3 H H sNp2 H 179 Ir 3 Pd Tn3 H H sSn1 H 180 Ir 3 Pd Tn3 H H sSn3 H 181 Ir 3 Pd Tn3 H H sPr H 182 Ir 3 Pd Tn3 H H sPr H 182 Ir 3 Pd Tn4 H sPr H 182 Ir 3 Pd Tn4 H sPr H 183 Ir 3 Pd Tn4 H sNp1 H 184 Ir 3 Pd Tn4		174	lr	3	Pd	Tn2	Н	Н	sPr	Н
177 Ir 3 Pd Tn3 H H sNp1 H 178 Ir 3 Pd Tn3 H H sNp2 H 179 Ir 3 Pd Tn3 H H sTn1 H 180 Ir 3 Pd Tn3 H H sPr H 181 Ir 3 Pd Tn3 H H sPr H 182 Ir 3 Pd Tn4 H H sPr H 183 Ir 3 Pd Tn4 H H sNp1 H 184 Ir 3 Pd Tn4 H H sNp2 H 185 Ir 3 Pd Tn4 H H sTn3 H 186 Ir 3 Pd Tn4 H H sTn3 H 187 Ir		175	lr	3	Pd	Tn2	Н	Н	sPe	Н
178 Ir 3 Pd Tn3 H H sNp2 H 179 Ir 3 Pd Tn3 H H sTn1 H 180 Ir 3 Pd Tn3 H H sFn3 H 181 Ir 3 Pd Tn3 H H sPr H 182 Ir 3 Pd Tn4 H H sPr H 183 Ir 3 Pd Tn4 H H sNp1 H 185 Ir 3 Pd Tn4 H H sNp2 H 186 Ir 3 Pd Tn4 H H sTn1 H 187 Ir 3 Pd Tn4 H H sTn1 H 189 Ir 3 Pd Tn4 H H sPr H 190 Ir		176	lr	3	Pd	Tn3	Н	Н	sPh	Н
179 Ir 3 Pd Tn3 H H sTn1 H 180 Ir 3 Pd Tn3 H H sTn3 H 181 Ir 3 Pd Tn3 H H sPP H 182 Ir 3 Pd Tn4 H H sPP H 183 Ir 3 Pd Tn4 H H sPP H 184 Ir 3 Pd Tn4 H H sNp1 H 185 Ir 3 Pd Tn4 H H sNp2 H 186 Ir 3 Pd Tn4 H H sTn3 H 187 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Tn4 H H sPr H 190 Ir <		177	lr	3	Pd	Tn3	Н	Н	sNp1	Н
180 Ir 3 Pd Tn3 H H sPr H 181 Ir 3 Pd Tn3 H H sPr H 182 Ir 3 Pd Tn3 H H sPr H 183 Ir 3 Pd Tn4 H H sPh H 184 Ir 3 Pd Tn4 H H sNp1 H 185 Ir 3 Pd Tn4 H H sNp2 H 186 Ir 3 Pd Tn4 H H sTn3 H 187 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Np1 H H sPp H 190 Ir <td< td=""><td></td><td>178</td><td>lr</td><td>3</td><td>Pd</td><td>Tn3</td><td>Н</td><td>Н</td><td>sNp2</td><td>Н</td></td<>		178	lr	3	Pd	Tn3	Н	Н	sNp2	Н
181 Ir 3 Pd Tn3 H H sPr H 182 Ir 3 Pd Tn3 H H sPr H 183 Ir 3 Pd Tn4 H H sPh H 184 Ir 3 Pd Tn4 H H sNp1 H 185 Ir 3 Pd Tn4 H H sNp2 H 186 Ir 3 Pd Tn4 H H sTn1 H 187 Ir 3 Pd Tn4 H H sTn3 H 188 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Tn4 H H sPr H 190 Ir 3 Pd Np1 H H sNp1 H 191 Ir <		179	lr	3	Pd	Tn3	Н	Н	sTn1	Н
182 Ir 3 Pd Tn3 H H sPe H 183 Ir 3 Pd Tn4 H H sPh H 184 Ir 3 Pd Tn4 H H sNp1 H 185 Ir 3 Pd Tn4 H H sNp2 H 186 Ir 3 Pd Tn4 H H sTn1 H 187 Ir 3 Pd Tn4 H H sTn3 H 188 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Tn4 H H sPr H 190 Ir 3 Pd Np1 H H sPp H 191 Ir 3 Pd Np1 H H sNp1 H 192 Ir <		180	ir	3	Pd	Tn3	I	I	sTn3	H
183 Ir 3 Pd Tn4 H H SPh H 184 Ir 3 Pd Tn4 H H SNp1 H 185 Ir 3 Pd Tn4 H H SNp2 H 186 Ir 3 Pd Tn4 H H sTn1 H 187 Ir 3 Pd Tn4 H H sTn3 H 188 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Tn4 H H sPr H 190 Ir 3 Pd Np1 H H sPr H 190 Ir 3 Pd Np1 H H sNp2 H 191 Ir 3 Pd Np1 H H sTn3 H 192 Ir		181	Ir	3	Pd	Tn3	I	Τ	sPr	Н
184 Ir 3 Pd Tn4 H H sNp1 H 185 Ir 3 Pd Tn4 H H sNp2 H 186 Ir 3 Pd Tn4 H H sTn1 H 187 Ir 3 Pd Tn4 H H sPr H 188 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Tn4 H H sPr H 190 Ir 3 Pd Np1 H H sPp H 190 Ir 3 Pd Np1 H H sNp1 H 191 Ir 3 Pd Np1 H H sNp2 H 192 Ir 3 Pd Np1 H H sPr H 194 Ir <		182	Ir	3	Pd	Tn3	I	I	sPe	Н
185 Ir 3 Pd Tn4 H H sNp2 H 186 Ir 3 Pd Tn4 H H sTn1 H 187 Ir 3 Pd Tn4 H H sSn3 H 188 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Tn4 H H sPe H 190 Ir 3 Pd Np1 H H sPe H 191 Ir 3 Pd Np1 H H sNp1 H 192 Ir 3 Pd Np1 H H sNp2 H 193 Ir 3 Pd Np1 H H sTn1 H 194 Ir 3 Pd Np1 H H sTn3 H 195 Ir		183	lr	3	₽d	Tn4	I	Н	sPh	Н
186 Ir 3 Pd Tn4 H H sTn1 H 187 Ir 3 Pd Tn4 H H sTn3 H 188 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Tn4 H H sPr H 190 Ir 3 Pd Np1 H H sPp H 190 Ir 3 Pd Np1 H H sNp1 H 191 Ir 3 Pd Np1 H H sNp2 H 192 Ir 3 Pd Np1 H H sNp2 H 193 Ir 3 Pd Np1 H H sTn3 H 194 Ir 3 Pd Np1 H H sPr H 195 Ir		184	lr	3	Pd	Tn4	Ι	Η	sNp1	Н
187 Ir 3 Pd Tn4 H H sTn3 H 188 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Tn4 H H sPr H 190 Ir 3 Pd Np1 H H sPp H 191 Ir 3 Pd Np1 H H sNp1 H 192 Ir 3 Pd Np1 H H sNp2 H 193 Ir 3 Pd Np1 H H sTn1 H 193 Ir 3 Pd Np1 H H sTn1 H 194 Ir 3 Pd Np1 H H sTn3 H 195 Ir 3 Pd Np1 H H sPr H 196 Ir		185	lr	3	Pd	Tn4	Н	Н	sNp2	Н
188 Ir 3 Pd Tn4 H H sPr H 189 Ir 3 Pd Tn4 H H sPe H 190 Ir 3 Pd Np1 H H sPh H 191 Ir 3 Pd Np1 H H sNp1 H 192 Ir 3 Pd Np1 H H sNp2 H 193 Ir 3 Pd Np1 H H sTn1 H 194 Ir 3 Pd Np1 H H sTn1 H 195 Ir 3 Pd Np1 H H sPr H 196 Ir 3 Pd Np1 H H sPr H 197 Ir 3 Pd Np2 H H sNp1 H 198 Ir <		186	lr	3	Pd	Tn4	H	Н	sTn1	н
189 Ir 3 Pd Tn4 H H sPe H 190 Ir 3 Pd Np1 H H sPh H 191 Ir 3 Pd Np1 H H sNp1 H 192 Ir 3 Pd Np1 H H sNp2 H 193 Ir 3 Pd Np1 H H sTn1 H 194 Ir 3 Pd Np1 H H sTn3 H 195 Ir 3 Pd Np1 H H sPr H 196 Ir 3 Pd Np1 H H sPr H 197 Ir 3 Pd Np2 H H sPh H 198 Ir 3 Pd Np2 H H sNp1 H 199 Ir <		187	lr	3	Pd	Tn4	Н	Н	sTn3	Н
190 Ir 3 Pd Np1 H H SPh H 191 Ir 3 Pd Np1 H H SNp1 H 192 Ir 3 Pd Np1 H H SNp2 H 193 Ir 3 Pd Np1 H H STn1 H 194 Ir 3 Pd Np1 H H STn3 H 195 Ir 3 Pd Np1 H H SPr H 196 Ir 3 Pd Np1 H H SPe H 197 Ir 3 Pd Np2 H H SNp1 H 198 Ir 3 Pd Np2 H H SNp2 H 199 Ir 3 Pd Np2 H H STn1 H 200 Ir 3 Pd Np2 H H STn1 H 201 Ir 3 Pd Np2 H H STn3 H 202 Ir 3 Pd Np2 H H SPr H 203 Ir 3 Pd Np2 H H SPr H		188	Ir	3	Pd	Tn4	Η	Н	sPr	Н
191 Ir 3 Pd Np1 H H SNp1 H 192 Ir 3 Pd Np1 H H SNp2 H 193 Ir 3 Pd Np1 H H STn1 H 194 Ir 3 Pd Np1 H H STn3 H 195 Ir 3 Pd Np1 H H SPr H 196 Ir 3 Pd Np1 H H SPe H 197 Ir 3 Pd Np2 H H SNp1 H 199 Ir 3 Pd Np2 H H SNp1 H 200 Ir 3 Pd Np2 H H STn1 H 201 Ir 3 Pd Np2 H H STn3 H 202 Ir 3 Pd Np2 H H STn3 H 203 Ir 3 Pd Np2 H H STn3 H		189	Ir	3	Pd	Tn4	Н	Н	sPe	Н
192 Ir 3 Pd Np1 H H sNp2 H 193 Ir 3 Pd Np1 H H sTn1 H 194 Ir 3 Pd Np1 H H sTn3 H 195 Ir 3 Pd Np1 H H sPr H 196 Ir 3 Pd Np1 H H sPe H 197 Ir 3 Pd Np2 H H sPh H 198 Ir 3 Pd Np2 H H sNp1 H 199 Ir 3 Pd Np2 H H sNp2 H 200 Ir 3 Pd Np2 H H sTn3 H 201 Ir 3 Pd Np2 H H sTn3 H 202 Ir		190	Ir	3	Pd	Np1	Н	Н	sPh	Н
193 Ir 3 Pd Np1 H H sTn1 H 194 Ir 3 Pd Np1 H H sTn3 H 195 Ir 3 Pd Np1 H H sPr H 196 Ir 3 Pd Np1 H H sPe H 197 Ir 3 Pd Np2 H H sPh H 198 Ir 3 Pd Np2 H H sNp1 H 199 Ir 3 Pd Np2 H H sNp2 H 200 Ir 3 Pd Np2 H H sTn1 H 201 Ir 3 Pd Np2 H H sTn3 H 202 Ir 3 Pd Np2 H H sPr H 203 Ir 3 Pd Np2 H H sPe H		191	lr	3	Pdi	Np1	Н	Н	sNp1	Н
194 Ir 3 Pd Np1 H H sTn3 H 195 Ir 3 Pd Np1 H H sPr H 196 Ir 3 Pd Np1 H H sPe H 197 Ir 3 Pd Np2 H H sPh H 198 Ir 3 Pd Np2 H H sNp1 H 199 Ir 3 Pd Np2 H H sNp2 H 200 Ir 3 Pd Np2 H H sTn3 H 201 Ir 3 Pd Np2 H H sTn3 H 202 Ir 3 Pd Np2 H H sPr H 203 Ir 3 Pd Np2 H H sPe H		192	lr	3	Pd	Np1	Η	Н	sNp2	Н
195 Ir 3 Pd Np1 H H sPr H 196 Ir 3 Pd Np1 H H sPe H 197 Ir 3 Pd Np2 H H sPh H 198 Ir 3 Pd Np2 H H sNp1 H 199 Ir 3 Pd Np2 H H sNp2 H 200 Ir 3 Pd Np2 H H sTn1 H 201 Ir 3 Pd Np2 H H sTn3 H 202 Ir 3 Pd Np2 H H sPr H 203 Ir 3 Pd Np2 H H sPe H		193	lr	3	Pđ	Np1	Н	Н	sTn1	Н
196 Ir 3 Pd Np1 H H sPe H 197 Ir 3 Pd Np2 H H sPh H 198 Ir 3 Pd Np2 H H sNp1 H 199 Ir 3 Pd Np2 H H sNp2 H 200 Ir 3 Pd Np2 H H sTn1 H 201 Ir 3 Pd Np2 H H sTn3 H 202 Ir 3 Pd Np2 H H sPr H 203 Ir 3 Pd Np2 H H sPe H		194	Ir	3	Pd	Np1	Н	Н	sTn3	Н
197 Ir 3 Pd Np2 H H sPh H 198 Ir 3 Pd Np2 H H sNp1 H 199 Ir 3 Pd Np2 H H sNp2 H 200 Ir 3 Pd Np2 H H sTn1 H 201 Ir 3 Pd Np2 H H sTn3 H 202 Ir 3 Pd Np2 H H sPr H 203 Ir 3 Pd Np2 H H sPe H		195	Ir	3	Pđ	Np1	Н	Н	sPr	Н
198 Ir 3 Pd Np2 H H sNp1 H 199 Ir 3 Pd Np2 H H sNp2 H 200 Ir 3 Pd Np2 H H sTn1 H 201 Ir 3 Pd Np2 H H sTn3 H 202 Ir 3 Pd Np2 H H sPr H 203 Ir 3 Pd Np2 H H sPe H		196	lr	3	Pd	Np1	Ι	Н	sPe	Η
199 Ir 3 Pd Np2 H H sNp2 H 200 Ir 3 Pd Np2 H H sTn1 H 201 Ir 3 Pd Np2 H H sTn3 H 202 Ir 3 Pd Np2 H H sPr H 203 Ir 3 Pd Np2 H H sPr H		197	lr	3	Pd	Np2	Ι	Н	sPh	Н
200 Ir 3 Pd Np2 H H sTn1 H 201 Ir 3 Pd Np2 H H sTn3 H 202 Ir 3 Pd Np2 H H sPr H 203 Ir 3 Pd Np2 H H sPe H		198	ir	3	Pd	Np2	Ι	Н	sNp1	Η
201 Ir 3 Pd Np2 H H sTn3 H 202 Ir 3 Pd Np2 H H sPr H 203 Ir 3 Pd Np2 H H sPe H		199	lr	3	Pd	Np2	Ħ	Н	sNp2	Н
202 Ir 3 Pd Np2 H H sPr H 203 Ir 3 Pd Np2 H H sPe H		200	ir	3	Pd	Np2	Н	Н	sTn1	Н
203 Ir 3 Pd Np2 H H sPe H		201	Ir	3	Pd	Np2	Н	Н	sTn3	Н
		202	Ir	3	Pd	Np2	Н	Н	sPr	Н
204 Ir 3 Pd Pe H H sPh H		203	lr	3	Pd	Np2	Н	Н	sPe	Н
	_	204	lr	3	Pd	Pe	н	Н	sPh	Н

Table 4 (continued)

No	Μ	m	CyN1	CyC1	R1	R2	R3	R4
205	lr	3	Pd	Pe	Н	Н	sNp1	Н
206	lr	3	Pd	Pe	Н	Н	sNp2	Н
207	lr	3	Pd	Pe	Н	Н	sTn1	Н
208	lr	3	Pd	Pe	Н	Н	sTn3	Н

Table 5

				Table 5				
No	М	m	CyN1	CyC1	R1	R2	R3	R4
209	Ir	3	Pd	Pe	Н	Ι	sPr	Н
210	lr	3	Pd	Pe	Η	Ι	sPe	Η
211	lr	3	Pd	Cn1	H	Ι	sPh	Н
212	lr	3	Pd	Cn1	Н	Ι	sNp1	Н
213	lr	3	Pd	Cn1	Н	Ι	sNp2	Н
214	lr	3	Pd	Cn1	I	Ι	sTn1	Н
215	lr	3	Pd	Cn1	Н	Н	sTn3	Н
216	lr	3	Pd	Cn1	Н	Τ	sPr	Н
217	lr	3	Pd	Cn1	Н	I	sPe	Ι
218	lr	3	Pd	Cn2	Н	Н	sPh	Η
219	Ir	3	Pd	Cn2	Н	Н	sNp1	Н
220	lr	3	Pd	Cn2	Н	Н	sNp2	Н
221	lr	3	Pd	Cn2	Н	Н	sTn1	Н
222	Ir	3	Pd	Cn2	Н	Н	sTn3	Η
223	lr	3	Pd	Cn2	Н	Н	sPr	Η
224	lr	3	Pd	Cn2	Н	Н	sPe	H
225	łr	3	Pd	Cz	Н	Н	sPh	Н
226	Ir	3	Pd	Cz	Н	Н	sNp1	Н
227	lr	3	Pd	Cz	Н	Н	sNp2	Н
228	Ir	3	Pd	Cz	Н	Н	sTn1	Н
229	lr	3	Pd	Cz	Н	Н	sTn3	Н
230	lr	3	Pd	Cz	Н	н	sPr	Н
231	lr	3	Pd	Cz	Н	н	sPe	Н
232	Ir	3	Pz	Ph	Н	Н	sPh	Н
233	Ir	3	Pz	Ph	Н	Н	sNp1	Н
234	lr	3	Pz	Ph	Н	Н	sNp2	Н
235	Ir	3	Pz	Ph	Н	Н	sTn1	Н
236	lr	3	Pz	Ph	Н	Н	sTn3	н
237	lr	3	Pz	Ph	Н	Н	sPr	Н
238	lr	3	Pz	Ph	Н	Н	sPe	Н
239	ir	3	Pz	Tn1	Н	Н	sPh	Н

Table 5 (continued)

No	М	m	CyN1	CyC1	R1	R2	R3	R4
240	lr	3	Pz	Tn1	Н	H	sNp1	Н
241	ir	3	Pz	Tn1	Н	Н	sNp2	Н
242	lr	3	Pz	Tn1	Н	Н	sTn1	Н
243	lr	3	Pz	Tn1	Н	Н	sTn3	Н
244	lr	3	Pz	Tn1	Н	Н	sPr	Н
245	lr	3	Pz	Tn1	Н	Н	sPe	Н
246	ir	3	Pz	Tn2	Н	Н	sPh	Н
247	Ir	3	Pz	Tn2	Н	Н	sNp1	Н
248	lr	3	Pz	Tn2	Н	Η	sNp2	Н
249	lr	з	Pz	Tn2	Н	Ŧ	sTn1	Н
250	lr	з	Pz	Tn2	Н	Н	sTn3	H
251	lr	თ	Pz	Tn2	I	Ι	sPr	Н
252	lr	თ	Pz	Tn2	Н	Ι	sPe	Н
253	lr	3	Pz	Tn3	Η	Ι	sPh	Н
254	lr	3	Pz	Tn3	I	I	sNp1	Н
255	lr	3	Pz	Tn3	Н	Н	sNp2	Н
256	lr	3	Pz	Tn3	Н	Н	sTn1	Н
257	lr	3	Pz	Tn3	Н	Н	sTn3	I
258	lr	3	Pz	Tn3	H	Н	sPr	Н
259	lr	3	Pz	Tn3	Н	Н	sPe	Н
260	lr	3	Pz	Tn4	Н	Н	sPh	Н

Table 6

N1-		F	0.14	0.01		T ===	r	г
No	М	m	CyN1	CyC1	R1	R2	R3	R4
261	Ir	3	Pz	Tn4	Н	Н	sNp1	Η
262	lr	3	Pz	Tn4	Н	н	sNp2	Н
263	lr	3	Pz	Tn4	Н	Н	sTn1	Η
264	lr	3	Pz	Tn4	Н	Н	sTn3	Н
265	lr	3	Pz	Tn4	Н	Н	sPr	Н
266	ir	3	Pz	Tn4	Н	Н	sPe	Н
267	lr	3	Pz	Np1	Н	Н	sPh	Н
268	Ιr	3	Pz	Np1	Н	Н	sNp1	Н
269	lr	3	Pz	Np1	Н	Н	sNp2	н
270	Ir	3	Pz	Np1	Н	Н	sTn1	Н
271	lr	3	Pz	Np1	Н	Н	sTn3	H
272	lr	3	Pz	Np1	Н	Н	sPr	Н
273	lr	3	Pz	Np1	Н	Н	sPe	Н
274	lr	3	Pz	Np2	Н	Н	sPh	Н

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Table 6 (continued)

No	М	m	CyN1	CyC1	R1	R2	R3	R4
275	lr	3	Pz	Np2	Н	H	sNp1	Н
276	lr	3	Pz	Np2	H	<u>''</u> Н	sNp2	Н
277	lr	3	Pz	Np2	Н	<u>''</u> Н		Н
278	lr	3	Pz				sTn1	
				Np2	н	H	sTn3	Н
279	lr ,	3	Pz	Np2	Н	Н	sPr	н
280	lr .	3	Pz	Np2	Н	Н	sPe	Н
281	lr	3	Pz	Pe	Н	Н	sPh	Н
282	lr .	3	Pz	Pe	Н	H	sNp1	Н
283	lr .	3	Pz	Pe	Н	Н	sNp2	H
284	lr	3	Pz	Pe	Н	Н	sTn1	H
285	lr	3	Pz	Pe	Н	Н	sTn3	Н
286	lr	3	Pz	Pe	Н	Н	sPr	Н
287	lr	3	Pz	Pe	Н	Н	sPe	Н
288	lr	3	Pz	Cn1	H	Н	sPh	Н
289	lr	3	Pz	Cn1	Η	Н	sNp1	Н
290	lr	3	Pz	Cn1	Н	Н	sNp2	Н
291	lr	3	Pz	Cn1	Н	Н	sTn1	Н
292	lr	3	Pz	Cn1	Н	Н	sTn3	Н
293	lr	3	Pz	Cn1	Н	Η	sPr	Н
294	lr	3	Pz	Cn1	Н	Н	sPe	Н
295	lr	3	Pz	Cn2	Н	Н	sPh	Н
296	lr	3	Pz	Cn2	Н	Н	sNp1	Н
297	lr	3	Pz	Cn2	Н	Н	sNp2	Н
298	lr	3	Pz	Cn2	Н	Н	sTn1	Н
299	lr	3	Pz	Cn2	Н	Н	sTn3	Н
300	lr	3	Pz	Cn2	Н	Н	sPr	Н
301	lr	3	Pz	Cn2	Н	Н	sPe	Н
302	lr	3	Pz	Cz	Н	Н	sPh	Н
303	lr	3	Pz	Cz	Н	Н	sNp1	Н
304	lr	3	Pz	Cz	Н	Н	sNp2	Н
305	lr	3	Pz	Cz	Н	Н	sTn1	Н
306	lr	3	Pz	Cz	Н	Н	sTn3	Н
307	lr	3	Pz	Cz	Н	Н	sPr	Н
308	lr	3	Pz	Cz	Н	Н	sPe	н
309	lr	3	Py1	Ph	Н	Н	sPh	Н
310	lr	3	Py1	Ρ́h	Н	Н	sNp1	Н
311	lr	3	Py1	Ph	Н	Н	sTn1	Н
312	lr	3	Py1	Ph	Н	Н	sTn3	Н

Table 7

					Table 7				
	No	М	m	CyN1	CyC1	R1	R2	R3	R4
5	313	lr	3	Py1	Tn1	Н	Н	sPh	Н
	314	lr	3	Py1	Tn1	Н	Н	sNp1	Н
	315	lr	3	Py1	Tn1	Н	Н	sTn1	Н
	316	lr	3	Py1	Tn1	н	Н	sTn3	Н
10	317	lr	3	Py1	Tn3	Н	Н	sPh	Н
	318	Ir	3	Py1	Tn3	Н	Н	sNp1	Н
	319	lr	3	Py1	Tn3	н	Н	sTn1	Н
15	320	lr	3	Py1	Tn3	Н	Н	sTn3	Н
	321	lr	3	Py1	Tn4	Н	Н	sPh	Н
	322	lr	3	Py1	Tn4	н	Н	sNp1	Н
20	323	lr	3	Py1	Tn4	н	Н	sTn1	н
20	324	lr	3	Py1	Tn4	Н	Н	sTn3	Н
	325	Ir	3	Py1	Np2	Н	Н	sPh	Н
	326	lr	3	Py1	Np2	Н	Н	sNp1	Н
25	327	lr	3	Py1	Np2	Н	Н	sTn1	Н
	328	lr	3	Py1	Np2	Н	Н	sTn3	Н
	329	lr	3	Py2	Ph	Н	н	sPh	Н
30	330	lr	3	Py2	Ph	Н	Н	sNp1	Н
30	331	lr	3	Py2	Ph	Н	Н	sTn1	Н
	332	lr	3	Py2	Ph	н	Н	sTn3	Н
	333	lr	3	Py2	Tn1	Н	Н	sPh	н
35	334	lr	3	Py2	Tn1	Н	Н	sNp1	Н
	335	ir	3	Py2	Tn1	Н	Н	sTn1	Н
	336	lr	3	Py2	Tn1	Н	н	sTn3	Н
40	337	lr	3	Py2	Tn3	Н	н	sPh	Н
	338	lr	3	Py2	Tn3	Н	Н	sNp1	Н
	339	lr .	3	Py2	Tn3	Н	Н	sTn1	Н
	340	ir	3	Py2	Tn3	н	Н	sTn3	н
45	341	ir	3	Py2	Tn4	н	Н	sPh	Н
	342	Ir	3	Py2	Tn4	Н	Н	sNp1	Н
	343	Ir	3	Py2	Tn4	н	Н	sTn1	Н
50	344	lr	3	Py2	Tn4	н	Н	sTn3	Н
	345	lr	3	Py2	Np2	Н	Н	sPh	Н
	346	lr .	3	Py2	Np2	Н	Н	sNp1	Н
	347	lr .	3	Py2	Np2	н	н	sTn1	Н
55	348	lr	3	Py2	Np2	Н	Н	sTn3	н
	349	lr	3	Pr	Ph	sPh	Н	Н	Н

Table 7 (continued)

No	М	m	CyN1	CyC1	R1	R2	R3	R4
350	lr	3	Pr	Ph	sNp2	Н	н	Н
351	lr	3	Pr	Ph	sTn1	Н	Н	Н
352	lr	3	Pr	Ph	sTn3	Н	Н	Н
353	lr	3	Pr	Tn1	sPh	Н	Н	Ι
354	Ir	3	Pr	Tn1	sNp2	Н	Н	Н
355	lr	3	Pr	Tn1	sTn1	Н	Н	Н
356	lr	· 3	Pr	Tn1	sTn3	н	Н	Н
357	lr	3	Pr	Tn3	sPh	Н	Н	Н
358	lr	3	Pr	Tn3	sNp2	Н	Н	H
359	lr	3	Pr	Tn3	sTn1	H	Н	Н
360	lr	3	Pr	Tn3	sTn3	Н	Н	Н
361	lr	3	Pr	Np2	sPh	н	Н	Н
362	lr	3	Pr	Np2	sNp2	Η	Η	Η
363	lr	3⋅	Pr	Np2	sTn1	H	Η	Н
364	lr	3	Pr	Np2	sTn3	Н	Н	Н

Table 8

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No	Μ	m	CyN1	CyC1	R1	R2	R3	R4
365	lr	3	Pz	Ph	sPh	Н	Н	Н
366	lr	3	Pz	Ph	sNp2	Н	Н	Н
367	lr	3	Pz	Ph	sTn1	Н	Н	Н
368	lr	3	Pz	Ph	sTn3	Н	Н	Н
369	lr	3	Pz	Tn1	sPh	Н	Н	Н
370	lr	3	Pz	Tn1	sNp2	Н	Н	Н
371	lr	3	Pz	Tn1	sTn1	Н	Н	Н
372	lr	3	Pz	Tn1	sTn3	Н	Н	Η
373	lr	3	Pz	Tn3	sPh	н	Н	H
374	lr	3	Pz	Tn3	sNp2	Н	Н	Н
375	lr	3	Pz	Tn3	sTn1	Н	Н	Н
376	lr	3	Pz	Tn3	sTn3	Н	Н	Н
377	lr	3	Pz	Np2	sPh	Н	Н	Н
378	lr	3	Pz	Np2	sNp2	Н	Н	Н
379	lr	3	Pz	Np2	sTn1	Н	Н	Н
380	lr	3	Pz	Np2	sTn3	Н	Н	Н

Table 9

No	М	m	CyN1	CyC1	R1	R2	R3	R4	R5	R6
381	lr	3	Pr	Ph	sPh	Н	н .	Н	Н	-NO2

`30

Table 9 (continued)

No						la	able 9 (d	continu	ed)			
383 Ir 3 Pr Ph STN1 H H H -CF3 H		No	М	m	CyN1	CyC1	R1	R2	R3	R4	R5	R6
383 Ir 3 Pr Ph STN1 H H H -CF3 H		382	lr	3	Pr	Ph	sNp2	Н	-СН3	Н	Н	Н
10 385 Ir 3 Pr Tn1 SPh H H H -OCH ₃ H 386 Ir 3 Pr Tn1 SNp2 H H H H SPh 387 Ir 3 Pr Tn1 STn1 H H H H SPh 388 Ir 3 Pr Tn1 STn3 H H H H SPh 389 Ir 3 Pr Tn3 SPh H H H H OCH ₃ H 390 Ir 3 Pr Tn3 SNp2 H H H H OCH ₃ H 391 Ir 3 Pr Tn3 STn1 H H H H OCH ₃ H 392 Ir 3 Pr Tn3 STn1 H H H H OCH ₃ H 393 Ir 3 Pr Tn3 STn3 H H H H OCH ₃ H 394 Ir 3 Pr Np2 SPh H H H H SPh 395 Ir 3 Pr Np2 STn1 H H H H SPh 396 Ir 3 Pr Np2 STn1 H H H H SPh 397 Ir 3 Pz Ph SPh H H OCH ₃ H 398 Ir 3 Pz Ph SPh H H OCH ₃ H 399 Ir 3 Pz Ph STn1 H H H H OCH ₃ 400 Ir 3 Pz Ph STn3 H H H H OCH ₃ 401 Ir 3 Pz Tn1 SPh H C3H7 H H H 402 Ir 3 Pz Tn1 SPh H C3H7 H H H 403 Ir 3 Pz Tn1 STn3 H H H H H H 404 Ir 3 Pz Tn1 STn3 H H H H H H 405 Ir 3 Pz Tn3 SPh H H H H H OCH ₃ 406 Ir 3 Pz Tn3 SPh H H H H OCH ₃ 407 Ir 3 Pz Tn3 STn1 H H H H OCH ₃ 408 Ir 3 Pz Tn3 STn1 H H H H OCH ₃ 409 Ir 3 Pz Tn3 STn1 H H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H OCH ₃ 400 Ir 3 Pz Tn3 STn1 H H H OCH ₃ 400 I	5	383	lr	3	Pr	Ph	sTn1	Н	Н	Н	-CF3	Н
10 386 ir 3 Pr Tn1 sNp2 H H H H SPh 387 ir 3 Pr Tn1 sTn1 H H H H H CF3 388 ir 3 Pr Tn1 sTn3 H H H H SPh 389 ir 3 Pr Tn3 sPh H H H H OCH3 H 390 ir 3 Pr Tn3 sNp2 H H H H OCH3 H 391 ir 3 Pr Tn3 sTn1 H H H H OCH3 H 392 ir 3 Pr Tn3 sTn1 H H H H OCH3 H 393 ir 3 Pr Np2 sPh H H H OCH3 H 394 ir 3 Pr Np2 sNp2 H H H H SPh 395 ir 3 Pr Np2 sNp2 H H H H SPh 396 ir 3 Pr Np2 sTn1 H H H H SPh 397 ir 3 Pr Np2 sTn3 H H H H OCH3 398 ir 3 Pr Np2 sTn3 H H H H OCH3 399 ir 3 Pr Ph sNp2 H H OCH3 H 399 ir 3 Pr Ph sNp2 H H OCH3 H 399 ir 3 Pr Ph sTn1 H H H H OCH3 400 ir 3 Pr Tn1 sNp2 H H H H H 402 ir 3 Pr Tn1 sNp2 H H H H H 403 ir 3 Pr Tn1 sTn1 H H H H H 404 ir 3 Pr Tn3 sNp2 H H H H H 405 ir 3 Pr Tn3 sNp2 H H H H OCH3 406 ir 3 Pr Tn3 sNp2 H H H H OCH3 407 ir 3 Pr Tn3 sNp2 H H H H OCH3 408 ir 3 Pr Tn3 sNp2 H H H H OCH3 409 ir 3 Pr Tn3 sNp2 H H H H H OCH3 400 ir 3 Pr Tn3 sNp2 H H H H OCH3 400 ir 3 Pr Tn3 sNp2 H H H H OCH3 400 ir 3 Pr Tn3 sNp2 H H H H H OCH3 400 ir 3 Pr Tn3 sNp2 H SC3H7 H H H H OCH3 400 ir 3 Pr Tn3 sNp2 H SC3H7 H H H H OCH3 400 ir 3 Pr Tn3 sNp2 H SC3H7 H H H OCH3 401 Ir 3 Pr Tn3 sNp2 H SC3H7 H H H H OCH3 401 Ir 3 Pr Np2 sNp2 H SC3H7 H H H H OCH3 401 Ir 3 Pr Np2 sNp2 H SC3H7 H H H H OCH3 401		384	lr	3	Pr	Ph	sTn3	Н	Н	Н	Н	sPh
387 Ir 3 Pr Tn1 STn1 H H H H H GF3 388 Ir 3 Pr Tn1 STn3 H H H H H SPh 389 Ir 3 Pr Tn3 SPh H H H H GCH3 H 390 Ir 3 Pr Tn3 SNp2 H H H H GCH3 391 Ir 3 Pr Tn3 STn1 H H H H GCH3 392 Ir 3 Pr Tn3 STn3 H H H H GCH3 H 394 Ir 3 Pr Np2 SPh H H H GCH3 H 395 Ir 3 Pr Np2 SNp2 H H H H SPh 395 Ir 3 Pr Np2 STn1 H H H H GCH3 396 Ir 3 Pr Np2 STn3 H H H H GCH3 397 Ir 3 Pr Np2 STn3 H H H H GCH3 398 Ir 3 Pr Np2 STn3 H H H GCH3 H 399 Ir 3 Pr Ph SPh H H GCH3 H 399 Ir 3 Pr Ph STn1 H H H H GCH3 400 Ir 3 Pr Tn1 SNp2 H H H H H 402 Ir 3 Pr Tn1 SNp2 H H H H H 403 Ir 3 Pr Tn1 STn3 H H H H H 404 Ir 3 Pr Tn3 SNP1 H H H H GCH3 406 Ir 3 Pr Tn3 SNP2 H H GCH3 H H 408 Ir 3 Pr Tn3 STn3 H H H H GCH3 409 Ir 3 Pr Tn3 STn3 H H H H GCH3 409 Ir 3 Pr Tn3 SNP2 H GC3H7 H H H H 410 Ir 3 Pr Np2 SNP2 H GC3H7 H H H H 410 Ir 3 Pr Np2 SNP2 H GC3H7 H H H 410 Ir 3 Pr Np2 SNP2 H GC3H7 H H H H 410 Ir 3 Pr Np2 SNP2 H GC3H7 H H H 410 Ir 3 Pr Np2 SNP2 H GC3H7 H H H 411 H H H H GCH3 412 Tn3 Tn3 Tn3 H H H H H H GCH3 413 Tn3 Tn3 Tn3 H H H H H GCH3 414 Tn3 Tn3 Tn3 H H H H H GCH3 415 Tn3 Tn3 Tn3 H H H H H GCH3 416 Tn3 Tn3 Tn3 H H H H H H GCH3 417 Tn3 Tn3 Tn3 H H H H H H GCH3 418 Tn3 Tn3 Tn3 H H H H H H H H H		385	lr	3	Pr	Tn1	sPh	Н	Н	Н	-OCH ₃	Н
388 3	10	386	ir	3	Pr	Tn1	sNp2	Н	Н	Н	Н	sPh
389 Ir 3 Pr Tn3 SPh H H H -OCH ₃ H 390 Ir 3 Pr Tn3 SNp2 H H H H -OCH ₃ 391 Ir 3 Pr Tn3 STn1 H H H H -OCH ₃ 392 Ir 3 Pr Tn3 STn3 H H H H -OCH ₃ 393 Ir 3 Pr Np2 SPh H H H -OCH ₃ H 394 Ir 3 Pr Np2 SNp2 H H H H SPh 395 Ir 3 Pr Np2 STn1 H H H H SPh 396 Ir 3 Pr Np2 STn3 H H H H H -OCH ₃ 397 Ir 3 Pz Ph SPh H H -OCH ₃ H H 398 Ir 3 Pz Ph SNp2 H H -OCH ₃ H H 399 Ir 3 Pz Ph STn1 H H H H -OCH ₃ 400 Ir 3 Pz Ph STn3 H H H H H -OCH ₃ 401 Ir 3 Pz Tn1 SPh H -C3H7 H H H 402 Ir 3 Pz Tn1 STn3 H H H H H H 403 Ir 3 Pz Tn1 STn3 H H H H H H 404 Ir 3 Pz Tn3 SPh H H H H H H 405 Ir 3 Pz Tn3 SPh H H H H H -OCH ₃ 406 Ir 3 Pz Tn3 SPh H H H H H -OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H -OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H -OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H -OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H -OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H -OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H -OCH ₃ 410 Ir 3 Pz Tn3 STn3 H H H H H -OCH ₃ 410 Ir 3 Pz Tn3 STn3 H H H H H -OCH ₃ 410 Ir 3 Pz Np2 SPh H H H H H -OCH ₃ 410 Ir 3 Pz Np2 SPh H -C3H7 H H H -OCH ₃		387	İr	3	Pr	Tn1	sTn1	Н	Н	Н	н	-CF3
390 Ir 3 Pr Tn3 SNp2 H H H H -OCH3 391 Ir 3 Pr Tn3 STn1 H H H H -OCH3 392 Ir 3 Pr Tn3 STn3 H H H H -OCH3 H 393 Ir 3 Pr Np2 SPh H H H -OCH3 H 394 Ir 3 Pr Np2 SNp2 H H H H SPh 395 Ir 3 Pr Np2 STn1 H H H H SPh 396 Ir 3 Pr Np2 STn3 H H H H H -OCH3 397 Ir 3 Pz Ph SPh H H -OCH3 H H 398 Ir 3 Pz Ph SNp2 H H -OCH3 H H 399 Ir 3 Pz Ph STn1 H H H H H -OCH3 400 Ir 3 Pz Ph STn3 H H H H H H 402 Ir 3 Pz Tn1 SPh H -C3H7 H H H H 403 Ir 3 Pz Tn1 STn1 H H H H H H 404 Ir 3 Pz Tn3 SPh H H H H H H 405 Ir 3 Pz Tn3 SPh H H H H H H 406 Ir 3 Pz Tn3 SPh H H H H H 408 Ir 3 Pz Tn3 STn3 H H H H H H 409 Ir 3 Pz Tn3 STn3 H H H H H H 409 Ir 3 Pz Tn3 STn3 H H H H H H 409 Ir 3 Pz Tn3 STn3 H H H H H H 409 Ir 3 Pz Tn3 STn3 H H H H H H -OCH3 410 Ir 3 Pz Tn3 STn3 H H H H H H -OCH3 410 Ir 3 Pz Np2 SPh H H H H H -OCH3 410 Ir 3 Pz Np2 SPh H H H H H H -OCH3 410 Ir 3 Pz Np2 SPh H H H H H H -OCH3 410 Ir 3 Pz Np2 SPh H H H H H H -OCH3 410 Ir 3 Pz Np2 SPh H H H H H H -OCH3 410 Ir 3 Pz Np2 SPh H -C3H7 H H H H H -OCH3 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H H -OCH3 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H H -OCH3		388	lr	3	Pr	Tn1	sTn3	H	Н	Н	Н	sPh
390 Ir 3 Pr Tn3 SNp2 H H H H -OCH3 391 Ir 3 Pr Tn3 STn1 H H H H -OCH3 392 Ir 3 Pr Tn3 STn3 H H H H -OCH3 393 Ir 3 Pr Np2 SPh H H H -OCH3 H 394 Ir 3 Pr Np2 SNp2 H H H H SPh 395 Ir 3 Pr Np2 STn1 H H H H SPh 396 Ir 3 Pr Np2 STn3 H H H H H -OCH3 397 Ir 3 Pz Ph SPh H H -OCH3 H H 398 Ir 3 Pz Ph SNp2 H H H H H -OCH3 399 Ir 3 Pz Ph STn1 H H H H -OCH3 400 Ir 3 Pz Ph STn3 H H H H H -OCH3 401 Ir 3 Pz Tn1 SPh H -C3H7 H H H 402 Ir 3 Pz Tn1 STn1 H H H H H H 403 Ir 3 Pz Tn1 STn1 H H H H H H 404 Ir 3 Pz Tn1 STn3 H H H H H H 405 Ir 3 Pz Tn3 SPh H H H H H 406 Ir 3 Pz Tn3 STn1 H H H H H 407 Ir 3 Pz Tn3 STn1 H H H H H 408 Ir 3 Pz Tn3 STn1 H H H H H 409 Ir 3 Pz Tn3 STn3 H H H H H 409 Ir 3 Pz Tn3 STn3 H H H H H 409 Ir 3 Pz Tn3 STn3 H H H H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H 411 H H H H H 412 H H H H H H 413 H H H H H H H 414 H H H H H H H 415 H H H H H H H H 416 H H H H H H H H H	15	389	ļr	3	Pr	Tn3	sPh	Ι	Ι	Н	-OCH ₃	Н
392 Ir 3 Pr Tn3 STn3 H H H -OCH ₃ H 393 Ir 3 Pr Np2 SPh H H H H -OCH ₃ H 394 Ir 3 Pr Np2 SNp2 H H H H SPh 395 Ir 3 Pr Np2 STn1 H H H H SPh 396 Ir 3 Pr Np2 STn3 H H H H H OCH ₃ 397 Ir 3 Pz Ph SPh H H -OCH ₃ H H 398 Ir 3 Pz Ph SNp2 H H -OCH ₃ H H 399 Ir 3 Pz Ph STn1 H H H H H OCH ₃ 400 Ir 3 Pz Ph STn3 H H H H H OCH ₃ 401 Ir 3 Pz Tn1 SPh H -C3H7 H H H H 402 Ir 3 Pz Tn1 STn1 H H H H H H 403 Ir 3 Pz Tn1 STn1 H H H H H H 404 Ir 3 Pz Tn1 STn3 H H H H H H 405 Ir 3 Pz Tn3 SPh H H H H H 406 Ir 3 Pz Tn3 STn1 H H H H OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H OCH ₃ 409 Ir 3 Pz Np2 SPh H OCH ₃ H H H OCH ₃ 410 Ir 3 Pz Np2 SPh H H H H H OCH ₃ 410 Ir 3 Pz Np2 SPh H H H H H OCH ₃ 410 Ir 3 Pz Np2 SPh H H H H H OCH ₃ 410 Ir 3 Pz Np2 SPh H H H H H OCH ₃ 410 Ir 3 Pz Np2 SPh H OCH ₃ H H H OCH ₃	15	390	lr	3	Pr	Tn3	sNp2	Ι	Ι	Н	Н	-OCH ₃
393 Ir 3 Pr Np2 SPh H H H H OCH ₃ H 394 Ir 3 Pr Np2 SNp2 H H H H SPh 395 Ir 3 Pr Np2 STn1 H H H H SPh 396 Ir 3 Pr Np2 STn3 H H H H H OCH ₃ 397 Ir 3 Pz Ph SPh H H OCH ₃ H H 398 Ir 3 Pz Ph STn1 H H H H OCH ₃ 399 Ir 3 Pz Ph STn1 H H H H OCH ₃ 400 Ir 3 Pz Ph STn3 H H H H H OCH ₃ 401 Ir 3 Pz Tn1 SPh H OC3H7 H H H H 402 Ir 3 Pz Tn1 STn1 H H H H H H 403 Ir 3 Pz Tn1 STn1 H H H H H H 404 Ir 3 Pz Tn1 STn3 H H H H H H 405 Ir 3 Pz Tn3 SPh H H H H H OCH ₃ 406 Ir 3 Pz Tn3 SNp2 H H OCH ₃ H H 408 Ir 3 Pz Tn3 STn3 H H H H H OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H OCH ₃ 409 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H 410 Ir 3 Pz Np2 SNp2 H OC3H7 H H H 410 Ir 3 Pz Np2 SNp2 H		391	lr	3	Pr	Tn3	sTn1	Ξ	Ι	H	Н	-OCH ₃
394 Ir 3 Pr Np2 SNp2 H H H H SPh 395 Ir 3 Pr Np2 STn1 H H H H SPh 396 Ir 3 Pr Np2 STn3 H H H H H OCH ₃ 397 Ir 3 Pz Ph SPh H H OCH ₃ H H 398 Ir 3 Pz Ph SNp2 H H OCH ₃ H H 399 Ir 3 Pz Ph STn1 H H H H OCH ₃ 400 Ir 3 Pz Ph STn3 H H H H OCH ₃ 401 Ir 3 Pz Tn1 SPh H OC3H7 H H H H 402 Ir 3 Pz Tn1 STn1 H H H H H H 403 Ir 3 Pz Tn1 STn1 H H H H H H 404 Ir 3 Pz Tn1 STn3 H H H H H H 405 Ir 3 Pz Tn3 SPh H H H H H 406 Ir 3 Pz Tn3 SNp2 H H OCH ₃ H H 408 Ir 3 Pz Tn3 STn3 H H H H H OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H OCH ₃ 410 Ir 3 Pz Np2 SPh H H H H H H OCH ₃		392	lr	3	Pr	Tn3	sTn3	Ι	H	H	-OCH ₃	Н
395 Ir 3 Pr Np2 STn1 H H H H H SPh 396 Ir 3 Pr Np2 STn3 H H H H H OCH ₃ 397 Ir 3 Pz Ph SPh H H OCH ₃ H H 398 Ir 3 Pz Ph SNp2 H H OCH ₃ H H 399 Ir 3 Pz Ph STn1 H H H H OCH ₃ 400 Ir 3 Pz Ph STn3 H H H H OCH ₃ 401 Ir 3 Pz Tn1 SPh H C3H7 H H H H 402 Ir 3 Pz Tn1 SNp2 H H H H H H 403 Ir 3 Pz Tn1 STn1 H H H H H H 404 Ir 3 Pz Tn1 STn3 H H H H H H 405 Ir 3 Pz Tn3 SPh H H H H H 406 Ir 3 Pz Tn3 SNp2 H H OCH ₃ H H 408 Ir 3 Pz Tn3 STn3 H H H H H OCH ₃ 409 Ir 3 Pz Np2 SNp2 H C3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H C3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H C3H7 H H H H 411 H H H H H H H H OCH ₃ 410 Ir 3 Pz Np2 SNp2 H C3H7 H H H H 411 H H H H H H H H 412 Ir 3 Pz Np2 SNp2 H C3H7 H H H H 413 Tn3	20	393	lr .	3	Pr	Np2	sPh	Τ	Τ	Н	-OCH ₃	Н
396 Ir 3 Pr Np2 STn3 H H H H -OCH ₃ 397 Ir 3 Pz Ph SPh H H -OCH ₃ H H 398 Ir 3 Pz Ph SNp2 H H -OCH ₃ H H 399 Ir 3 Pz Ph STn1 H H H H -OCH ₃ 400 Ir 3 Pz Ph STn3 H H H H H 402 Ir 3 Pz Tn1 SPh H -C3H7 H H H 402 Ir 3 Pz Tn1 STn1 H H H H H 403 Ir 3 Pz Tn1 STn1 H H H H H 404 Ir 3 Pz Tn1 STn3 H H H H H 405 Ir 3 Pz Tn3 SPh H H H H H 406 Ir 3 Pz Tn3 SNp2 H H H H H 407 Ir 3 Pz Tn3 STn1 H H H -OCH ₃ 409 Ir 3 Pz Tn3 STn3 H H H H H 408 Ir 3 Pz Tn3 STn3 H H H H H 409 Ir 3 Pz Np2 SPh H H H H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H H 410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H H 411 H H H H H H H 412 H H H H H H H H 414 H H H H H H H H H		394	lr	3	Pr	Np2	sNp2	I	Ι	H	Н	sPh
397 Ir 3 Pz Ph sPh H H -OCH ₃ H H 398 Ir 3 Pz Ph sNp2 H H -OCH ₃ H H 399 Ir 3 Pz Ph sTn1 H H H H H -OCH ₃ 400 Ir 3 Pz Ph sTn3 H H H H H H 401 Ir 3 Pz Tn1 sPh H -C3H7 H H H 402 Ir 3 Pz Tn1 sNp2 H H H H H 403 Ir 3 Pz Tn1 sTn1 H H H H H 404 Ir 3 Pz Tn1 sTn3 H H H H H sPh 405 Ir 3 Pz Tn3 sPh H H H H H -OCH ₃ 406 Ir 3 Pz Tn3 sNp2 H H -OCH ₃ H H 408 Ir 3 Pz Tn3 sTn1 H H H H -OCH ₃ 409 Ir 3 Pz Np2 sPh H H H H H -OCH ₃ 410 Ir 3 Pz Np2 sNp2 H -C3H7 H H H H 410 Ir 3 Pz Np2 sNp2 H -C3H7 H H H H 410 Ir 3 Pz Np2 sNp2 H -C3H7 H H H H 4110 Ir 3 Pz Np2 sNp2 H -C3H7 H H H H 412 H H H H H H H H H		395	lr	3	Pr	Np2	sTn1	I	Ι	Н	Н	sPh
397	25	396	lr	3	Pr	Np2	sTn3	Ι	Τ	H	Н	-OCH₃
399 Ir 3 Pz Ph STn1 H H H H -OCH ₃ 400 Ir 3 Pz Ph STn3 H H H H H -OCH ₃ 401 Ir 3 Pz Tn1 SPh H -C3H7 H H H 402 Ir 3 Pz Tn1 SNp2 H H H H H H 403 Ir 3 Pz Tn1 STn1 H H H H H 404 Ir 3 Pz Tn1 STn3 H H H H H SPh 405 Ir 3 Pz Tn3 SPh H H H H H -OCH ₃ 406 Ir 3 Pz Tn3 SNp2 H H -OCH ₃ H H 407 Ir 3 Pz Tn3 STn1 H H H -OCH ₃ H H 408 Ir 3 Pz Tn3 STn1 H H H -OCH ₃ H H 409 Ir 3 Pz Np2 SPh H H H H H -OCH ₃	23	397	lr	3	Pz	Ph	sPh	Ξ	T	-OCH3	Н	Н
30		398	lr	3	Pz	Ph	sNp2	Τ	Ŧ	-OCH ₃	Н	Н
401 Ir 3 Pz Tn1 sPh H -C3H7 H H H H 402 Ir 3 Pz Tn1 sNp2 H H H H H H 403 Ir 3 Pz Tn1 sTn1 H H H H H 404 Ir 3 Pz Tn1 sTn3 H H H H H sPh 405 Ir 3 Pz Tn3 sPh H H H H H 406 Ir 3 Pz Tn3 sNp2 H H -OCH3 H H 407 Ir 3 Pz Tn3 sTn1 H H -OCH3 H H 408 Ir 3 Pz Tn3 sTn3 H H H H H -OCH3 409 Ir 3 Pz Np2 sPh H H H H H H 410 Ir 3 Pz Np2 sNp2 H -C3H7 H H H H 4110 Ir 3 Pz Np2 sNp2 H -C3H7 H H H H 401 H H H H H H H 402 H H H H H H H 403 H H H H H H H H 404 H H H H H H H 405 H H H H H H H H 406 H H H H H H H 407 H H H H H H H H 408 H H H H H H H H 409 H H H H H H H H H		399	lr	3	Pz	Ph	sTn1	H	Н	Н	Н	-OCH ₃
402 Ir 3 Pz Tn1 sNp2 H H H H H H H H H	30	400	lr	3	Pz	Ph	sTn3	Н	H	Н	Н	-OCH ₃
403 Ir 3 Pz Tn1 sTn1 H H H H H H H H SPh 404 Ir 3 Pz Tn1 sTn3 H H H H H sPh 405 Ir 3 Pz Tn3 sPh H H H H -OCH ₃ 406 Ir 3 Pz Tn3 sNp2 H H -OCH ₃ H H 407 Ir 3 Pz Tn3 sTn1 H H -OCH ₃ H H 408 Ir 3 Pz Tn3 sTn3 H H H H H -OCH ₃ 409 Ir 3 Pz Np2 sNp2 H -C3H7 H H H H 410 Ir 3 Pz Np2 sNp2 H -C3H7 H H H H 410 Ir 3 Pz Np2 sNp2 H -C3H7 H H H H 401 H H H H H H H 402 H H H H H H H 403 H H H H H H H 404 H H H H H H H 405 H H H H H H 406 H H H H H H 407 H H H H H H H 408 H H H H H H H 409 H H H H H H H 409 H H H H H H H H 400 H H H H H H H 400 H H H H H H H H H		401	lr	3	Pz	Tn1	sPh	Н	-C3H7	H	Н	Н
404 Ir 3 Pz Tn1 sTn3 H H H H sPh 405 Ir 3 Pz Tn3 sPh H H H H -OCH ₃ 406 Ir 3 Pz Tn3 sNp2 H H -OCH ₃ H H 40 407 Ir 3 Pz Tn3 sTn1 H H -OCH ₃ H H 408 Ir 3 Pz Tn3 sTn3 H H H H -OCH ₃ 409 Ir 3 Pz Np2 sNp2 H -C3H7 H H H 410 Ir 3 Pz Np2 sNp2 H -C3H7 H H H 411 H H H H 405 H SPh H H H H 406 H SPh H H H H 407 H SPh H H H H 408 H SPh H H H H 409 H SPh H H H H 409 H SPh H H H H 410 H SPh H SPh 410 H SPh H H H H 407 H SPh H H H 408 H SPh H H H H 409 H SPh H H H H 409 H SPh H H H H 400 H SPh H H H H H H 400 H SPh H H H H H H 400 H SPh H H H H H H 400 H SPh H H H H H H 400 H SPh H H H H H H 400 H SPh H H H H H H H 400 H SPh H H H H H H H 400 H SPh H H H H H H H H 400 H SPh H H H H H H H H 400 H SPh H H H H H H H H H 400 H SPh H H H H H H H H H		402	lr	3	Pz	Tn1	sNp2	Н	н	Н	Н	Н
405 Ir 3 Pz Tn3 sPh H H H H -OCH ₃ 406 Ir 3 Pz Tn3 sNp2 H H -OCH ₃ H H H H H H H H H	35	403	lr	3	Pz	Tn1	sTn1	Н	H	Н	Н	Н
406 ir 3 Pz Tn3 sNp2 H H -OCH ₃ H H 407 ir 3 Pz Tn3 sTn1 H H -OCH ₃ H H 408 ir 3 Pz Tn3 sTn3 H H H H -OCH ₃ 409 ir 3 Pz Np2 sPh H H H H -OCH ₃ 410 ir 3 Pz Np2 sNp2 H -C3H7 H H H		404	lr	3	Pz	Tn1	sTn3	Н	Н	H	Н	sPh
407 Ir 3 Pz Tn3 sTn1 H H -OCH ₃ H H H H H H H H H		405	ir	3	Pz	Tn3	sPh	Н	Н	H	Н	-OCH ₃
408 Ir 3 Pz Tn3 sTn3 H H H H -OCH ₃ 409 Ir 3 Pz Np2 sPh H H H H -OCH ₃ 410 Ir 3 Pz Np2 sNp2 H -C3H7 H H H		406	ir	3	Pz	Tn3	sNp2	Н	н	-OCH ₃	Н	Н
409 Ir 3 Pz Np2 sPh H H H H -OCH ₃ 410 Ir 3 Pz Np2 sNp2 H -C3H7 H H H	40	407	lr	3	Pz	Tn3	sTn1	Н	Н	-OCH₃	Н	Н
410 Ir 3 Pz Np2 sNp2 H -C3H7 H H H		408	lr	3	Pz	Tn3	sTn3	Н	н	Н	Н	-OCH ₃
410 Ir 3 Pz Np2 SNp2 H -C3H7 H H H		409	lr	3	Pz	Np2	sPh	Н	н	Н	н	-0CH ₃
	45	410	lr	3	Pz	Np2	sNp2	Н	-C3H7	Н	Н	н

Table 10

No	М	m	CyN1	CyC1	R1	R2	R3	R4	R5	R6
411	lr	3	Pz	Np2	sTn1	Н	н	-CF3	Н	Н
412	lr	3	Pz	Np2	sTn3	Н	Н	-CF3	Н	Н
413	lr	3	Та	Ph	C4H9	C4H9	sPh	Н	оснз	н
414	lr	3	Pr	Ph	sPh	Н	Н	Н	Н	Н
415	lr	3	Pr	Ph	sNp2	Н	-СН3	Н	Н	Н
416	ir	3	Pr	Ph	sTn1	Н	Н	н	Н	Н

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Table 10 (continued)

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No	M	m	CyN1	CyC1	R1	R2	R3	R4	R5	R6
417	lr	3	Pr	Ph	sTn3	Н	Н	Н	Τ	Н
418	lr	3	Pr	Tn1	sPh	Н	Н	Н	-OCH ₃	Н
419	lr	3	Pr	Tn1	sNp2	Н	Н	Н	Н	Н
420	lr	3	Pr	Tn1	sTn1	Н	Н	Н	Н	Н
421	lr	3	Pr	Tn1	sTn3	Н	H	Н	Н	Н
422	lr	3	Pr	Tn3	sPh	Н	Н	Н	-OCH ₃	Н
423	lr	3	₽r	Tn3	sNp2	Н	Н	Н	Н	Н
424	lr	3	Pr	Tn3	sTn1	Н	-NO2	Н	Н	Н
425	I r	3	Pr	Tn3	sTn3	Н	Н	Н	Н	Н
426	ir	3	Pr	Np2	sPh	н	н	Н	н	Н
427	Ir	3	Pr	Np2	sNp2	Н	Н	Н	Н	Н
428	lr	3	Pr	Np2	sTn1	н	Н	Н	Н	Н
429	lr	3	Pr	Np2	sTn3	Н	Н	Н	Н	Н
430	lr	3	Pz	Ph	sPh	Н	Н	-F ,	Н	Н
431	ŀ٢	3	Pz	Ph	sNp2	Н	Н	Н	Н	Н
432	ir	3	Pz	Ph	sTn1	-CN	Н	н	Н	Н
433	1r	3	Pz	Ph	sTn3	Н	Н	Н	Н	Н
434	lr	3	Pz	Tn1	sPh	H	-C3H7	Н	Н	Н
435	lr	3	Pz	Tn1	sNp2	Н	Η.	-CH2-CH=CH-CH3	Н	Н
436	lr	3	Pz	Tn1	sTn1	Н	н	Н	H	Н
437	lr	3	Pz	Tn1	sTn3	Н	Ξ	Н	Н	H
438	lr	3	Pz	Tn3	sPh	Н	-SC3H7	Н	Н	Н
439	lr	3	Pz	Tn3	sNp2	Н	Н	Н	Н	Н
440	lr	3	Pz	Tn3	sTn1	Н	Н	Н	Н	Н
441	lr	3	Pz	Tn3	sTn3	Н	Н		Н	Н
442	lr	3	Pz	Np2	sPh	Н	Н	Н	Н	Н
443	lr	3	Pz	Np2	sNp2	Н	Н	Н	Н	Н
444	lr	3	Pz	Np2	sTn1	Н	Н	Н	Н	Н
445	lr	3	Pz	Np2	sTn3	Н	Н	Н	Н	Н

Table 11

No	М	m	n	CyN1	CyC1	CyN2	CyC2	R1	R2	R3	R4	R1'	R2'	R3'	R4'
446	lr	2	1	Pr	Ph	Pr	Tn1	sPh	Н	Н	Н	sPh	Н	Н	Н
447	lr	2	1	Pr	Ph	Pr	Tn1	sNp2	Н	Н	Н	sNp2	Н	н	Н
448	lr	2	1	Pr	Ph	Pr	Tn1	sTn1	Н	н	Н	sTn1	Н	Н	Н
449	lr	2	1	Pr	Ph	Pr	Tn1	sTn3	Н	Н	Н	sTn3	Н	Н	Н
450	Ir	2	1	Pr	Tn3	Pr	Np2	sPh	Н	Н	Н	sPh	Н	Н	Н
451	Ir	2	1	Pr	Tn3	Pr	Np2	sNp2	Н	н	Н	sNp2	Н	Н	н

Table 11 (continued)

No	М	Э	n	CyN1	CyC1	CyN2	CyC2	R1	R2	R3	R4	R1'	R2'	R3'	R4'
452	lr	2	1	Pr	Tn3	Pr	Np2	sTn1	H	Н	Н	sTn1	Η	Н	Н
453	lr .	2	1	Pr	Tn3	Pr	Np2	sTn3	Н	Η	Н	sTn3	н	Н	Н

Table 12

					ıaı	ole 12					
No	М	m	n	CyN1	CyC1	E	G	R1	R2	R3	R4
454	lr	lr	1	Pr	Ph	-СН3	-СН3	sPh	Н	Н	Н
455	lr	lr	1	Pr	Ph	-СН3	-СН3	sNp2	Н	Н	Н
456	lr	lr	1	Pr	Ph	-СН3	-СН3	sTn1	Н	Н	Н
457	lr	lr	1	Pr	Ph	-СН3	-СН3	Н	Н	sTn3	Н
458	lr	lr	1	Pr	Tn3	-СН3	sPh	Н	Н	sPh	Н
459	lr	lr	1	Pr	Tn3	-СН3	sPh	Н	Н	sNp2	н
460	lr	lr	1	Pr	Tn3	-СН3	sPh	Н	Н	sTn1	Н
461	lr	lr	1	Pr	Tn3	-СН3	sPh	Н	Н	sTn3	Н

Table 13

				Table 13				
No	М	m	CyN1	CyC1	R1	R2	R3	R4
462	Rh	3	Pr	Ph	sPh	Н	Ι	Н
463	Rh	3	Pr	Ph	sNp2	Н	Н	Н
464	Rh	3	Pr	Ph	sTn1	Н	Н	Η
465	Rh	3	Pr	Ph	sTn3	Н	Н	Н
466	Rh	3	Pr	Tn1	sPh	Н	Η	Н
467	Rh	3	Pr	Tn1	sNp2	Н	Н	Н
468	Rh	3	Pr	Tn1	sTn1	Н	Н	Н
469	Rh	3	Pr	Tn1	sTn3	Н	Η	Н
470	Rh	3	Pr	Tn3	sPh	Н	H	Н
471	Rh	3	Pr	Tn3	sNp2	Н	Η	Н
472	Rh	3	Pr	Tn3	sTn1	Н	H.	Η
473	Rh	3	Pr	Tn3	sTn3	Н	Τ	Η
474	Rh	3	Pr	Np2	sPh	Н	Η	Н
475	Rh	3	Pr	Np2	sNp2	Н	Н	Н
476	Rh	3	Pr	Np2	sTn1	Н	H	Н
477	Rh	3	Pr	Np2	sTn3	Н	Н	Н

Table 14

No	Δ	m	CyN1	CyC1	R1	R2	R3	R4
478	Pt	2	Pr	Ph	sPh	Н	Н	Н .
479	Pt	2	Pr	Ph	sNp2	Н	Н	Н

Table 14 (continued)

No	М	m	CyN1	CyC1	R1	R2	R3	R4
480	Pt	2	Pr	Ph	sTn1	н	H	Η
481	Pt	2	Pr	Ph	sTn3	Н	Н	Η
482	Pt	2	Pr	Tn1	sPh	Н	Н	Н
483	Pt	2	Pr	Tn1	sNp2	Н	Н	Н
484	Pt	2	Pr	Tn1	sTn1	Н	Н	Н
485	Pt	2	Pr	Tn1	sTn3	Н	Н	Н
486	Pt	2	Pr	Tn3	sPh	H.	Н	Н
487	Pt	2	Pr	Tn3	sNp2	Η	Н	Н
488	Pt	2	Pr	Tn3	sTn1	Η	Н	Н
489	Pt	2	Pr	Tn3	sTn3	Н	Н	Н
490	Pt	2	Pr	Np2	sPh	Н	Н	Н
491	Pt	2	Pr	Np2	sNp2	Н	Н	Н
492	Pt	2	Pr	Np2	sTn1	Н	Н	Н
493	Pt	2	Pr	Np2	sTn3	Н	Н	H

Table 15

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Table 15								
No	М	m	CyN1	CyC1	R1	R2	R3	R4
494	Pd	2	Pr	Ph	sPh	Н	Н	Η
495	Pd	2	Pr	Ph	sNp2	Н	Η	Н
496	Pd	2	Pr	Ph	sTn1	Н	Н	Н
497	Pd	2	Pr	Ph	sTn3	Н	Н	Н
498	Pd	2	Pr	Tn1	sPh	Н	Н	Н
499	Pd	2	Pr	Tn1	sNp2	Н	Η	Н
500	Pd	2	Pr	Tn1	sTn1	Н	Ι	Η
501	Pd	2	Pr	Tn1	sTn3	Н	Η	Н
502	Pd	2	Pr	Tn3	sPh	Н	Н	H
503	Pd	2	Pr	Tn3	sNp2	Н	Н	Η
504	Pd	2	Pr	Tn3	sTn1	H	I	Η
505	Pd	2	Pr	Tn3	sTn3	Н	Τ	Ι
506	Pd	2	Pr	Np2	sPh	Н	Η	Н
507	Pd	2	Pr	Np2	sNp2	Н	Н	Н
508	Pd	2	Pr	Np2	sTn1	н	Н	Н
509	Pd	2	Pr	Np2	sTn3	Н	Н	Н

Table 16

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No	М	m	CyN1	CyC1	R1	R2	R3	R4	R5	R6
510	lr	က	Pr	Ph	sPe	H	Н	Н	Н	H
511	İr	3	Pr	Ph	sPh	Н	sPh_	Н	~	H
512	lr	3	Pr	Ph	Н	-0	sPh	н	Н	~>
513	Ir	3	Pr	Np2	sPe	Н	Н	Н	Н	Н
514	lr	3	Pr	Np2	Н	Н	sTn1	H	CH3	I
515	lr	3	Pr	Tn1	CH3	Н	sTn1	Н	CH3	Н
516	lr	3	Pr	Tn1	sPh	Н	sTn1	Н	sPh	Ξ

Table 17

No	М	m	ก	CyN1	CyC1	R1	R2	R3	R4	E	G
517	lr	2	1	Pr	Tn3	н	I	sPh	Н	СНЗ	СНЗ
518	lr	2	1	Pr	Tn1	Н	Н	sTn1	Н	СНЗ	СНЗ
519	lr	2	1	Pr	Np2	Н	Н	sNp2	Н	СНЗ	СНЗ
520	lr	3	0	Py1	Ph	sPh	Н	Н	Н	-	-
521	lr	3	0	Py1	Ph	sNp1	Н	Н	Н	-	-
522	lr	3	0	Pr	Ph	Н	Н	Н	sPh	-	-
523	lr	3	0	Pr	Ph	Н	sPh	Н	Н	-	-
524	lr	3	0	Pr	Tn1	Ph	Н	Н	Н	-	-
525	lr	2	1	Py1	Ph	sPh	Н	Н	Н	СНЗ	СНЗ
526	lr	2	1	Py1	Ph	sNp1	Н	H	Н	СНЗ	СНЗ
527	lr	2	1	Pr	Ph	Н	Н	Н	sPh	СНЗ	СНЗ
528	lr	2	1	Pr	Ph	Н	sPh	Н	Н	СНЗ	СНЗ
529	lr	2	1	Pr	Tn1	Ph	Н	Н	Н	СНЗ	СНЗ

[0059] Hereinbelow, the present invention will be described more specifically based on Examples.

Examples 1 - 6

[0060] Each of luminescence devices having a layer structure shown in Figure 1B were prepared in the following

[0061] On a 1.1 mm-thick glass substrate (transparent substrate 15), a 100 nm-thick film (transparent electrode 14) of ITO (indium tin oxide) was formed by sputtering, followed by patterning to form a stripe electrode including 100 lines each having a width of 100 nm and a spacing with an adjacent line of 10 nm (i.e., electrode pitch of 110 nm).

[0062] On the ITO-formed substrate, three organic layers and two metal electrode layers shown below were successively formed by vacuum (vapor) deposition using resistance heating in a vacuum chamber (10⁻⁴ Pa).

Organic layer 1 (hole transport layer 13) (40 nm): α-NPD

Organic layer 2 (luminescence layer 12) (30 nm): co-deposited film of CBP:metal complex (metal coordination compound shown in Table 20) (95:5 by weight)

Organic layer 3 (electron transport layer 16) (30 nm): Alq3

Metal electrode layer 1 (metal electrode 11) (15 nm): Al-Li alloy (Li = 1.8 wt. %)

Metal electrode layer 2 (metal electrode 11) (100 nm): Al

[0063] The above-deposited metal electrode layers 1 and 2 (Al-Li layer and Al layer) had a stripe electrode pattern including 100 lines each having a width of 100 nm and a spacing of 10 nm (electrode pitch = 110 nm) and arranged so that the stripe electrode pattern intersected with that of the ITO electrode at right angles to form a matrix of pixels each having an effective electrode area of 3 mm² comprising 20 ITO lines bundled together at a lead-out portion and

15 Al (Al-Li) lines bundled together at a lead-out portion.

[0064] Each of the thus-prepared luminescence devices was taken out of the vacuum chamber and was subjected to a continuous energization (current passage) test in an atmosphere of dry nitrogen gas stream so as to remove device deterioration factors, such as oxygen and moisture (water content).

[0065] The continuous energization test was performed by continuously applying a voltage at a constant current density of 50 mA/cm² to the luminescence device having the ITO (transparent) electrode (as an anode) and the AI (metal) electrode (as a cathode), followed by measurement of emission luminance (brightness) with time so as to determine a time (luminance half-life) required for decreasing an initial luminance (60 - 220 cd/m²) to 1/2 thereof.

[0066] The results are shown in Table 18 appearing hereinafter.

Comparative Example 1

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[0067] A comparative luminescence device was prepared and evaluated in the same manner as in Examples 1 - 6 except that the Ir complexes (metal coordination compounds shown in Table 20) was changed to Ir-phenylpyrimidine complex (Ir(ppy)₃) shown below.

[0068] The results are also shown in Table 18 below.

Table 18

Ex. No.	Compound No.	Luminance half-life (Hr)
Ex. 1	3	450
Ex. 2	11	550
Ex. 3	22	500
Ex. 4	43	500
Ex. 5	45	600
Ex. 6	385	400
Ex. 7	413	650
Comp.Ex. 1	Ir(ppy) ₃	300

[0069] As is apparent from Table 18, compared with the conventional luminescence device using Ir(ppy)₃, the luminescence devices using the metal coordination compounds of formula (1) according to the present invention provide longer luminance half-lives, thus resulting in an EL device having a high durability (luminance stability) based on a good stability of the metal coordination compound of formula (1) of the present invention.

Example 7

[0070] A color organic EL display apparatus shown in Figure 2 was prepared in the following manner.

[0071] An active matrix substrate had a planar structure basically similar to a structure described in U.S. Patent No. 6,114,715.

[0072] Specifically, on a 1.1 mm-thick glass substrate, top state-type TFTs of polycrystalline silicon were formed in an ordinary manner and thereon, a flattening film was formed with contact holes for electrical connection with a pixel electrode (anode) at respective source regions, thus preparing an active matrix substrate with a TFT circuit.

[0073] On the active matrix substrate, a 700 nm-thick pixel electrode (anode) of ITO having a larger work function

was formed in a prescribed pattern. On the ITO electrode, prescribed organic layers and a 100 nm-thick Al electrode (cathode) were successively formed by vacuum deposition with a hard mask, followed by patterning to form a matrix of color pixels (128x128 pixels).

[0074] The respective organic layers corresponding to three color pixels (red (R) green (G) and blue (B)) were consisting of the following layers.

<R pixel region>

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[0075] α-NPD (40 nm)/CBP: Ex. Comp. No. 22 (93:7 by weight) (30 nm)/BCP (20 nm)/Alq 3 (40 nm)

<G pixel region>

[0076] α -NPD (50 nm)/Alq 3 (50 nm)

15 <B pixel region>

[0077] α -NPD (50 nm)/BCP (20 nm)/Alq 3 (50 nm)

[0078] When the thus-prepared color organic EL display apparatus was driven, desired color image data can be displayed stably with good image qualities.

Example 8 (Synthesis of Example Compound No. 22)

[0079]

[0080] In a 500 ml-three-necked flask, 12.6 g (85.2 mM) of 2,5-dichloropyridine, 15.2 g (85.4 mM) of benzothiophene-2-boronic acid, 75 ml of toluene, 37.5 ml of ethanol and 75 ml of 2M-sodium carbonate aqueous solution were placed and stirred at room temperature under nitrogen stream, and 3.06 g (2.64 mM) of tetrakis(triphenylphosphine)palladium (0) was added thereto, followed by refluxing under stirring for 8 hours under nitrogen stream. After the reaction, the reaction mixture was cooled on an ice bath to precipitate a crystal, which was then filtered out and washed with water. To the crystal, 100 ml of methanol was added and washed under stirring at room temperature, followed by filtration to recover the crystal. The crystal was purified by silica gel column chromatography (eluent: chloroform) and recrystallized from a mixture solvent of chloroform-methanol to obtain 11.8 g (Yield: 56.4 %) of 5-chloro-2-(benzo[b]thienyl)pyridine (colorless crystal).

[0081] In a 100 ml-three-necked flask, 4.91 g (20.0 mM) of 5-chloro-2-(benzo[b]thienyl)pyridine, 3.66 g (30.0 mM) of phenylboronic acid, 9.58 g (40.0 mM) of tripotassium phosphate hydrate, 3.2 mg (0.020 mM) of palladium (II) acetate, 11.9 mg (0.040 mM) of 2-di-tert-butylphosphinobiphenyl and 60 ml of toluene were placed and refluxed under stirring for 24 hours at 100 °C under nitrogen stream. After the reaction, the reaction mixture was cooled on an ice bath to precipitate a crystal, which was then filtered out and washed with water. To the crystal, 25 ml of methanol was added and washed under stirring at room temperature, followed by recovery by filtration. The crystal was purified by silica gel column chromatography (eluent: chloroform) and recrystallized from a chloroform-methanol mixture solvent to obtain 1.17 g (Yield: 20.4 %) of 2-(benzo[b]thienyl)-5-phenylpyridine (colorless crystal).

[0082] In a 100 ml-four-necked flask, 50 ml of glycerol was placed and heated at 130 - 140 °C under stirring and bubbling with nitrogen for 2 hours. Then, the glycerol was cooled by standing to 100 °C, and 1.15 g (4.00 mM) of 2-(benzo[b]lhienyl)-5-phenylpyridine and 0.40 g (0.82 mM) of iridium (III) acetylacetonate were added thereto, followed by sturing for 5 hours at 180 - 235 °C under nitrogen stream. The reaction mixture was cooled to room temperature and poured into 300 ml of 1N-hydrochloric acid to form a precipitate. The precipitate was recovered by filtration and washed with water, followed by drying for 5 hours at 100 °C under reduced pressure. The resultant precipitate was silica gel column chromatography (eluent: chloroform) to obtain 0.26 g (Yield: 30.2 %) of red powdery tris[2-(benzo[b] thicnyl)-5-phenylpyridine-C²,N]iridium (III).

[0083] According to MALDI-TOF MS (matrix-assisted laser desorption ionization-time of flight mass spectroscopy), the compound exhibited M+ (mass number of the corresponding cation formed by removal of 1 electron) of 1051.2, thus confirming the objective iridium complex.

[0084] When the compound was dissolved in toluene and subjected to measurement of phosphorescence spectrum at an excited light wavelength of 380 nm by using a fluorescence spectrometer, the compound exhibited a phosphorescence spectrum showing \$\lambda\$max\$ (maximum emission wavelength) of 620 nm, thus confirming clear red luminescence. [0085] When the luminescence device prepared in 5 Example 3 using the above-synthesized metal coordination compound (Ex. Comp. No. 22) was subjected to measurement of phosphorescence spectrum in a similar manner, a clear red luminescence was confirmed similarly as in the case of the compound in toluene 0 described above.

Example 9 (Synthesis of Ex. Comp. No. 11)

[0086] A metal coordination compound (Ex. Comp. No. 11) was synthesized through the following reaction schemes. Hereinafter, the synthesis yield is simply represented by "Y".

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Y. 10 %

[0087] According to MALDI-TOF MS, the compound exhibited M+ = 919.0, thus being identified as the objective iridium compound.

[0088] When the compound was dissolved in toluene and subjected to measurement of phosphorescence spectrum at an excited light wavelength of 400 nm by using a fluorescence spectrometer, the compound exhibited a phosphorescence spectrum showing \(\text{\text{max}} \) (maximum emission wavelength) of 612 nm, thus confirming clear red luminescence.

[0089] When a luminescence device having a layer structure shown below and using the above-synthesized metal coordination compound (Ex. Comp. No. 11) was prepared and subjected to measurement of phosphorescence spectrum in a similar manner, a clear red luminescence was confirmed similarly as in the case of the compound in toluene described above.

[0090] ITO (100 nm)/ α -NPD (40 nm)/CBP: Ex. Comp. No. 11 (95:5 by weight)(30 nm)/BCP (20 nm)/Alq3 (40 nm)/Al-Li (1 nm)/Al (100 nm).

[0091] Further, the luminescence device exhibited a good rectifying characteristic.

[0092] Specifically, Figure 3A is a graph showing a relationship between an electric field strength (E) and a current density of the luminescence device, and Figure 3B is a graph showing a relationship between an electric field strength (E) and a luminance (L) of the luminescence device. Further, Figure 3C shows a luminescence spectrum of the luminescence device under application of a voltage of 10 volts.

[0093] The luminescence device exhibited a luminescence efficiency of 0.8 1m/W under application of a voltage of 10 volts. The luminescence device also emitted stable luminescence even when the luminescence device was continuously supplied with the voltage for ca. 200 hours.

35 Example 10 (Synthesis of Ex. Comp. No. 45)

[0094] A metal coordination compound (Ex. Comp. No. 45) was synthesized through the following reaction schemes.

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[0095] According to MALDI-TOF MS, the compound exhibited $M^+ = 1183.3$, thus being identified as the objective iridium compound.

[0096] When the compound was dissolved in toluene and subjected to measurement of phosphorescence spectrum at an excited light wavelength of 380 nm by using a fluorescence spectrometer, the compound exhibited a phosphorescence spectrum showing λ max (maximum emission wavelength) of 603 nm, thus confirming clear reddish orange luminescence.

[0097] When the luminescence device prepared in Example 5 using the above-synthesized metal coordination compound (Ex. Comp. No. 45) was subjected to measurement of phosphorescence spectrum in a similar manner, a clear reddish orange luminescence was confirmed similarly as in the case of the compound in toluene described above.

[0098] Further, the luminescence device exhibited a good rectifying characteristic.

[0099] The luminescence device exhibited a luminescence efficiency of 0.5 lm/W under application of a voltage of 8 volts. The luminescence device also emitted stable luminescence even when the luminescence device was continuously supplied with the voltage for ca. 150 hours.

Example 11 (Another synthesis of Ex. Comp. No. 22)

[0100] Tris[2-(benzo[b]thienyl)-5-phenylpyridine-C²,N]iridium (III) (Ex. Comp. No. 22) prepared in Example 8 was synthesized through another reaction schemes shown below.

[0101] In a 200 ml-three-necked flask, 0.58 mg (1.64 mmole) of iridium (III) chloride-trihydrate (made by Across Organics Co.), 1.5 g (5.22 mmole) of 2-(benzo[b]thienyl)-5-phenylpyridine, 45 ml of ethoxyethanol and 15 ml of water were placed and stirred for 30 min. at room temperature under nitrogen stream, followed by 24 hours of reflux under stirring. The reaction product was cooled to room temperature, and the precipitate was recovered by filtration and washed with water, followed successive washing with ethanol and acetone. After drying under a reduced pressure at room temperature, 1.02 g of red powdery tetrakis[2-(benzo[b]thienyl)-5-phenylpyridine-C²,N]-(µ-dichloro)diiridium (III)

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was obtained.

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[0102] In a 200 ml-three-necked flask, 70 ml of ethoxyethanol, 0.95 g (0.72 mmole) of tetrakis[2-(benzo[b]thienyl)-5-phenylpyridine- C^2 ,N](μ -dichloro)-diiridium (III), 0.22 g (2.10 mM) of acetylacetone and 1.04 g (9.91 mM) of sodium carbonate, were placed and stirred for 1 hour at room temperature under nitrogen stream and then refluxed under stirring for 15 hours. The reaction product was cooled with ice, and the precipitate was filtered out and washed with water. The precipitate was then purified by silica gel column chromatography (eluent: chloroform/methanol = 30/1) to obtain 0.43 g of red powdery bis[2-(benzo[b]thienyl)-5-phenylpyridine- C^2 ,N](acetylacetonato)-iridium (III) (Example Compound No. 517). According to MALDI-TOF MS, M+ of 864.2 of the compound was confirmed. A toluene solution of the compound exhibited a luminescence spectrum showing λ max = 631 nm and a quantum yield of 0.18 relative to 1.0 of Ir(ppy)₃.

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$$CH_3$$
 CH_3 CH_3 CH_3 CH_3

[0103] In a 100 ml-three-necked flask, 0.27 g (0.94 mM) of 2-(benzo[b]thienyl)-5-phenylpyridine, 0.36 g (0.42 mM) of bis[2-benzo[b]thienyl)-5-phenylpyridine-C²,N](acetylacetonato)iridium (III) and 25 ml of glycerol, were placed and heated around 180 °C for 8 hours under stirring and nitrogen stream. The reaction product was cooled to room temperature and poured into 170 ml of 1N-hydrochloric acid, and the precipitate was filtered out, washed with water and dried at 100 °C under a reduced pressure for 5 hours. The precipitate was purified by silica gel column chromatography with chloroform as the eluent to obtain 0.27 g of red powdery tris[2-(benzo[g]thienyl-5-phenylpyridine-C²,N]iridium (III) (Example Compound No. 22). According to MALDI-TOF MS, M⁺ of 1051.2 of the compound was confirmed. A toluene solution of the compound exhibited a luminescence spectrum showing λmax = 627 nm and a quantum yield of 0.17

relative to 1.0 of Ir(ppy)₃.

[0104] The above-synthesized compound and a luminescence device prepared by using the compound exhibited luminescence characteristics similar to those of the compound and luminescence device prepared in Example 8.

[0105] Bis[2-(benzo[g]thienyl)-5-phenylpyridine-C²,N]iridium (III) (Ex. Comp. No. 517) prepared in this example as an intermediate product exhibited λmax which was longer by ca. 4 nm than that of the final product (Ex. Comp. No. 22) having three identical ligands. Further, when a luminescence device using the intermediate product was prepared and evaluated in the same manner as in Example 8, the luminescence device exhibited a luminescence spectrum showing λmax = 631 nm. Accordingly, the intermediate product used in this example can also be used as a luminescence material.

Example 12 (Another synthesis of Ex. Comp. No. 45)

[0106] The metal coordination compound (Ex. Comp. No. 45) prepared in Example 10 was synthesized through another reaction schemes shown below.

$$\frac{2 \times \operatorname{IrCl_{3}.3H_{2}O}}{(1)}$$

[0107] In a 200 ml-three-necked flask, 0.58 mg (1.64 mmole) of iridium (III) chloride-trihydrate (made by Across Organics Co.), 1.7 g (5.1 mmole) of a compound (1), 45 ml of ethoxyethanol and 15 ml of water were placed and stirred for 30 min. at room temperature under nitrogen stream, followed by 24 hours of reflux under stirring. The reaction product was cooled to room temperature, and the precipitate was recovered by filtration and washed with water, followed successive washing with ethanol and acetone. After drying under a reduced pressure at room temperature, 1.0 g (yield = 93.4 %) of red powdery compound (2) was obtained.

[0108] In a 200 ml-three-necked flask, 70 ml of ethoxyethanol, 0.90 g (0.71 mmole) of the compound (2), 0.22 g (2.10 mmole) of acetylacetone and 1.04 g (9.91 mmole) of sodium carbonate, were placed and stirred for 1 hour at room temperature under nitrogen stream and then refluxed under stirring for 15 hours. The reaction product was cooled with ice, and the precipitate was filtered out and washed with water. The precipitate was then purified by silica gel column chromatography (eluent: chloroform/methanol = 30/1) to obtain 0.39 g of red powdery compound (3) (Example Compound No. 519). According to MALDI-TOF MS, M+ of 952.3 of the compound was confirmed. A toluene solution of the compound exhibited a luminescence spectrum showing λmax = 608 nm and a higher quantum yield of 0.30 relative to 1.0 of lr(ppy)₃ in this emission wavelength region.

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$$CH_3$$

$$CH_3$$

$$(1)$$

$$(4)$$

[0109] In a 100 ml-three-necked flask, 0.29 g (0.88 mM) of the compound (1) 0.34 g (0.35 mM) of the compound (3) and 25 ml of glycerol, were placed and heated around 180 °C for 8 hours under stirring and nitrogen stream. The reaction product was cooled to room temperature and poured into 170 ml of 1N-hydrochloric acid, and the precipitate was filtered out, washed with water and dried at 100 °C under a reduced pressure for 5 hours. The precipitate was purified by silica gel column chromatography with chloroform as the eluent to obtain 0.23 g of red powdery compound
 (4) (Example Compound No. 45). According to MALDI-TOF MS, M+ of 1183.4 of the compound was confirmed. A toluene solution of the compound exhibited a luminescence spectrum showing λmax = 603 nm and a quantum yield of 0.278 relative to 1.0 of Ir(ppy)₃.

[0110] The above-synthesized compound and a luminescence device prepared by using the compound exhibited luminescence characteristics similar to those of the compound and luminescence device prepared in Example 10. [0111] The compound (3) (Ex. Comp. No. 519) prepared in this example as an intermediate product exhibited \(\lambda\) max which was longer by ca. 4 nm than that of the final product (Ex. Comp. No. 45) having three identical ligands. Further, when a luminescence device using the intermediate product was prepared and evaluated in the same manner as in

Example 10, the luminescence device exhibited a luminescence spectrum showing λmax = 608 nm and an external luminescence yield of 0.7 lm/W. Further, the luminescence device emitted stable luminescence even when continuously supplied with the voltage for ca. 100 hours. Accordingly, the intermediate product used in this example can also be used as a luminescence material.

Example 13 (Synthesis of Ex. Comp. Nos. 520 and 525)

- 40 [0112] It is easy to synthesize the following compounds in the same manner as in Example 11 except that 4-chloropyrimidine is synthesized from 4(3H)-pyrimidone (made by Aldrich Co.) in the same manner as the process described at pages 37 and 38 of JP-A (Tokuhyo) 2001-504113 (corr. to U.S. Patent No. 6,300,330) and is reacted with 4-phenyl-boronic acid (made by Lancaster Co.) to obtain 4-(biphenyl-4-yl)pyrimidine, which is used instead of 2-(benzo[b]thienyl)-5-phenylpyridine.
- 45 [0113] Bis[4-(biphenyl-4-yl)pyridine-C³,N³] (acetylacetonato) iridium (III) (Ex. Comp. No. 520).
 [0114] Tris[4-(biphenyl-4-yl)pyrimidine-C³,N³] iridium (III) (Ex. Comp. No. 525).

Example 14 (Synthesis of Ex. Comp. Nos. 521 and 526)

- [0115] It is easy to synthesize the following compounds in the same manner as in Example 11 except that 4-(4-chlorophenyl)pyrimidine is synthesized from 4-chloropyrimidine prepared in Example 13 and 4-chlorophenylboronic acid (made by Aldrich Co.) and was reacted with 2-naphthaleneboronic acid (made by Lancaster Co.) to obtain 4-[4-(2-naphthyl)phenyl]-pyrimidine, which is used instead of 2-(benzo[b]thienyl)-5-phenylpyridine.
 - [0116] Bis{4-[4-(2-naphthyl)phenyl]pyrimidine-C³,N³}(acetylacetonato)iridium (III) (Ex. Comp. No. 521).
- 55 [0117] Tris{4-[4-(2-naphthyl)phenyl]pyrimidine-C³,N³}iridium (III) (Ex. Comp. No. 526).

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Example 15 (Synthesis of Ex. Comp. Nos. 522 and 527)

[0118] It is easy to synthesize the following compounds in the same manner as in Example 11 except that 2,4-diphenylpyridine is synthesized from phenylboronic acid (made by Tokyo Kasei Kogyo K.K.) and 4-phenyl-2-bromopyridine (made by General Intermediates of Canada) and was used instead of 2-(benzo[b]thienyl)-5-phenylpyridine.

[0119] Bis(2,4-diphenylpyridine-C²,N¹)(acetylacetonato)iridium (III) (Ex. Comp. No. 522).

[0120] Tris(2,4-diphenylpyridine-C²,N¹)iridium (III) (Ex. Comp. No. 527).

Example 16 (Synthesis of Ex. Comp. Nos. 523 and 528)

[0121] It is easy to synthesize the following compounds in the same manner as in Example 11 except that 2-(biphenyl-3-yl)pyridine is synthesized from 3-biphenylboronic acid (made by Lancaster Co.) and 2-bromopyridine (made by Tokyo Kasei Kogyo K.K.) and is used instead of 2-(benzo[b]thienyl)-5-phenylpyridine.

[0122] Bis[2-(biphenyl-3-yl)pyridine-C⁴,N³)(acetylacetonato)iridium (III) (Ex. Comp. No. 523).

[0123] Tris[2-(biphenyl-2-yl)pyridine-C⁴,N³)iridium (III) (Ex. Comp. No. 528).

Example 17 (Synthesis of Ex. Comp. Nos. 524 and 529)

[0124] It is easy to synthesize the following compounds in the same manner as in Example 11 except that 2-(5-bro-mothiophene-2-yl)pyridine is synthesized from 2-bromopyridine (made by Tokyo Kasei Kogyo K.K.) and 5-bromothiophene-2-boronic acid (made by Aldrich Co.) and was reacted with phenylboronic acid (made by Tokyo Kasei Kogyo K.K.) to obtain 2-(5-phenylthiophene-2-yl)pyridine, which is used instead of 2-(benzo[b]thienyl)-5-phenylpyridine.

[0125] Bis[2-(5-phenylthiophene-2-yl)pyridine-C2,N1)(acetylacetonato)iridium (III) (Ex. Comp. No. 524).

[0126] Tris[2-(5-phenylthiophene-2-yl)pyridine-C2,N1)iridium (III) (Ex. Comp. No. 529).

[0127] As described above, according to the present invention, the metal coordination compound of the formula (1) characterized by aromatic substituent. The electroluminescence device (luminescence device) of the present invention using, as a luminescent center material, the metal coordination compound of the formula (1) is an excellent device which not only allows high-efficiency luminescence but also retains a high luminance for a long period and shows little deterioration by current passage. Further, the display apparatus using the electroluminescence device of the present invention exhibits excellent display performances.

[0128] An electroluminescence device having a layer containing a specific metal coordination compound is provided. The metal coordination compound is represented by formula (1) below:

$$ML_mL_n'$$
 (1),

wherein M is a metal atom of Ir, Pt, Rh or Pd; L and L' are mutually different bidentate ligands; m is 1, 2 or 3 and n is 0, 1 or 2 with the proviso that m+n is 2 or 3; a partial structure MLm is represented by formula (2) shown below and a partial structure ML'_n is represented by formula (3) or (4) shown below:

The metal coordination compound of the formula (1) is characterized by having at least one aromatic substituent for at least one of CyN1, CyN2, CyC1 and CyC2. The metal coordination compound having the aromatic substituent is effective in providing high-efficiency luminescence, long-term high luminance, and less deterioration by current passing.

55 Claims

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1. A metal coordination compound represented by formula (1) below:

$$ML_mL_n'$$
 (1),

wherein M is a metal atom of Ir, Pt, Rh or Pd; L and L' are mutually different bidentate ligands; m is 1, 2 or 3 and n is 0, 1 or 2 with the proviso that m+n is 2 or 3; a partial structure MLm is represented by formula (2) shown below and a partial structure ML'_n is represented by formula (3) or (4) shown below:

wherein CyN1 and CyN2 are each cyclic group capable of having a substituent, including a nitrogen atom and bonded to the metal atom M via the nitrogen atom; CyC1 and CyC2 are each cyclic group capable of having a substituent, including a carbon atom and bonded to the metal atom M via the carbon atom with the proviso that the cyclic group CyN1 and the cyclic group CyC1 are bonded to each other via a covalent bond and the cyclic group CyN2 and the cyclic group CyC2 are bonded to each other via a covalent bond;

the optional substituent of the cyclic groups is selected from a halogen atom, cyano group, a nitro group, a trialkylsilyl group of which the alkyl groups are independently a linear or branched alkyl group having 1 to 8 carbon atoms, a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -O-CO-, -CH=CH-or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom; or an aromatic group capable of having a substituent which is selected from an aromatic group capable of having a substituent (that is a halogen atom, a cyano atom, a nitro atom, a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -O-CO-, -CH=CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom), a halogen atom, a cyano atom, a nitro atom, and a linear or branched alkyl group having 1 to 20 carbon atoms (of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -O-CO-, -O-CO-, -CH=CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom);

E and G are independently a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom, or an aromatic group capable of having a substituent (that is a halogen atom, a cyano atom, a nitro atom, a trialkylsilyl group of which the alkyl groups are independently a linear or branched alkyl group having 1 - 8 carbon atoms, a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -C-CO-, -CH-CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom; and

the cyclic groups CyN1, CyN2, CyC1 and CyC2 have at least one aromatic substituent capable of having a substituent which is selected from an aromatic group capable of having a substituent (that is a halogen atom, a cyano atom, a nitro atom, a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -O-CO-, -CH=CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom), a halogen atom, a cyano atom, a nitro atom, a linear or branched alkyl group having 1 to 20 carbon atoms of which the alkyl group can include one or non-neighboring two or more methylene groups that can be replaced with -O-, -S-, -CO-, -CO-O-, -O-CO-, -CH=CH- or -C=C-, and the alkyl group can include a hydrogen atom that can be optionally replaced with a fluorine atom).

- A metal coordination compound according to Claim 1, including a partial structure ML'_n represented by the formula
 (3) in the formula (1).
- A metal coordination compound according to Claim 1, including a partial structure ML'_n represented by the formula (4) in the formula (1).

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- 4. A metal coordination compound according to Claim 1, wherein n is 0 in the formula (1).
- 5. A metal coordination compound according to Claim 1, wherein in the formula (2), CyN1 is pyridyl group and CyC1 is naphthyl group.
- 6. A metal coordination compound according to Claim 1, wherein in the formula (2), CyN1 is pyridyl group and CyC1 is thienyl group.
- 7. A metal coordination compound according-to Claim 1, wherein in the formula (2), CyN1 is pyridyl group and CyC1 is benzothienyl group.
 - 8. An electroluminescence device, comprising: a pair of electrodes disposed on a substrate, and a luminescence unit comprising at least one organic compound disposed between the electrodes, wherein the organic compound comprises a metal coordination compound represented by the formula (1) in Claim 1.
 - 9. A metal coordination compound according to Claim 8, including a partial structure ML'_n represented by the formula (3) in the formula (1).
 - A metal coordination compound according to Claim 8, including a partial structure ML'_n represented by the formula (4) in the formula (1).
 - 11. A metal coordination compound according to Claim 8, wherein n is 0 in the formula (1).
- 12. An electroluminescence device according to Claim 8, wherein a voltage is applied between the electrodes to emit light.
 - 13. An electroluminescence device according to Claim 8, wherein a voltage is applied between the electrodes to emit phosphorescence.
- 30 14. A picture display apparatus, comprising an electroluminescence device according to Claim 8, and a means: for supplying electric signals to the electroluminescence device.

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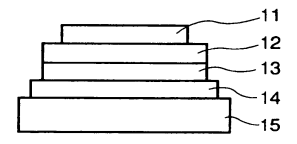


FIG. 1A

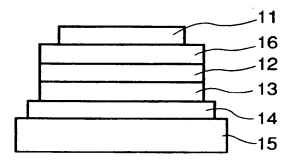


FIG. 1B

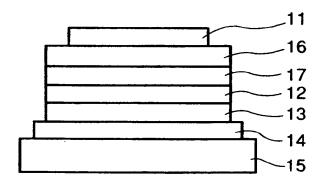


FIG. 1C

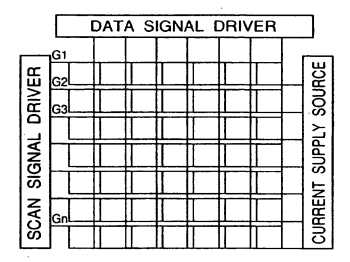


FIG. 2

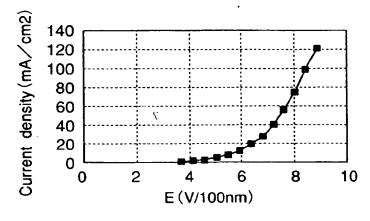


FIG. 3A

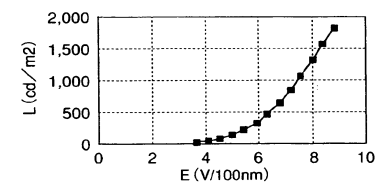


FIG. 3B

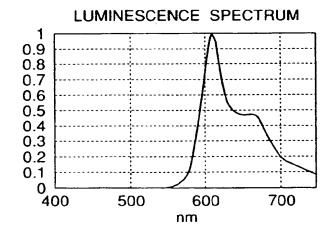


FIG. 3C

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EUROPEAN PATENT APPLICATION

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- (54)Metal coordination compound, luminescene device and display apparatus
- (57)An electroluminescence device having a layer containing a specific metal coordination compound is provided. The metal coordination compound is represented by formula (1) below:

$$ML_mL_n'$$
 (1),

wherein M is a metal atom of Ir, Pt, Rh or Pd; L and L' are mutually different bidentate ligands; m is 1, 2 or 3 and n is 0, 1 or 2 with the proviso that m+n is 2 or 3; a partial structure MLm is represented by formula (2) shown below and a partial structure ML'_n is represented by formula (3) or (4) shown below:

$$M = \begin{cases} CyN1 \\ CyC1 \end{cases} m \qquad (2) \qquad M = \begin{cases} CyN2 \\ CyC2 \end{cases} n \qquad (3) \qquad M = \begin{cases} O = E \\ O = G \end{cases} n \qquad (4)$$

The metal coordination compound of the formula (1) is characterized by having at least one aromatic substituent for at least one of CyN1, CyN2, CyC1 and CyC2. The metal coordination compound having the aromatic substituent is effective in providing high-efficiency luminescence, long-term high luminance, and less deterioration by current passing.



Application Number EP 02 00 5113

	DOCUMENTS CONSID	ERED TO BE RELE	VANT		
Category	Citation of document with it of relevant pass.	ndication, where appropriate ages	,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CI.7)
E	EP 1 191 612 A (CAM 27 March 2002 (2002 * the whole documer	2-03-27)		,2,4 - 9, 1-14	H01L51/30 C07F15/00 H01L51/00
E	EP 1 191 613 A (CAM 27 March 2002 (2002 * the whole documer	?-03-27)		,2,4,8, 9,11-14	
P,X	EP 1 175 128 A (FUS 23 January 2002 (20 * the whole documen	02-01-23))]	-4,8-14	·
X	DJUROVICH P I ET AL CYCLOMETALATED COMP PHOSPHORESCENT EMIT AND ORGANIC LEDS" POLYMER PREPRINTS, SOCIETY, US, vol. 41, no. 1, Mar	PLEXES ÀS EFFICIE TERS IN POLYMER AMERICAN CHEMICA Ch 2000 (2000-03	NT BLEND	,2,4,8,),11-14	
	pages 770-771, XP00 ISSN: 0032-3934 * the whole documer				TECHNICAL FIELDS SEARCHED (Int.Ct.7)
P,X	WO 02/02714 A (PETR PONT (US); WANG YIN VLADIMI) 10 January * the whole documer	IG (US); GRUSHIN / 2002 (2002-01-1	-	4,8-14	H01L
P,X	WO 02/15645 A (UNIV SOUTHERN CALIFORNIA DISPLAY C) 21 Febru * the whole documen	n (US); UNIVÉRSAL Jary 2002 (2002-0	i	4,8-14	
E	EP 1 211 257 A (CAM 5 June 2002 (2002-0 * the whole documen	06-05)	1	14	
	•	-/-	-		
	The present search report has	been drawn up for all claims			
	Place of search	Date of completion of	the search		Examiner
	Munich	3 Februar		Кое	ssler, J-L
X : parti Y : parti docu A : tech	ATEGORY OF CITED DOCUMENTS icularly relevant if taken alone icularly relevant if combined with anot iment of the same category nological background	E : ear afte her D : doc L : doc	ory or principle un lier patent docum r the filing date sument cited in th ument cited for o	ent, but publis e application ther reasons	hed on, or
O : non- P : inter	-written disclosure rmediate document	8 : me	mber of the same ument	patent family,	corresponding



Application Number

EP 02 00 5113

Category	Citation of document with indi-	cation, where appropriate.	Relevant	CLASSIFICATION OF THE
Category	of relevant passage		to claim	APPLICATION (Int.CI.7)
E .	WO 02/45466 A (CANON (JP); KAMATANI JUN (6 June 2002 (2002-06 * the whole document	JP); MIURA SEISHI (J) -06)	1-14	
Ē	WO 03/000661 A (MIZU SATOSHI (JP); CANON I 3 January 2003 (2003 * the whole document	KK (JP); FURUGÓRI MAN) -01-03)	1-14	
P,X	WO 01/041512 A (UNIV SOUTHERN CALIFORNIA 7 June 2001 (2001-06 * the whole document	(US)) -07)	1-14	
X	ON ELECTROPHOSPHORES APPLIED PHYSICS LETTI INSTITUTE OF PHYSICS	EMITTING DEVICES BASED CENCE" ERS, AMERICAN . NEW YORK, US, IY 1999 (1999-07-05),	1,2,4,8, 9,11-14	TECHNICAL FIELDS SEARCHED (Int.Cl.7)
X	THE PREPARATION OF A PHOTOREDUCING AGENTS TRIS-ORTHO-METALATED	SERIES OF STRONG: FAC COMPLEXES OF IRIDIUM ED 2-PHENYLPYRIDINES" AMERICAN CHEMICAL L, pages 1685-1687,	1-4,8-14	
	The present search report has bee	en drawn up for all claims		
	Place of search	Date of completion of the search	<u> </u>	Examiner
	Munich	3 February 2004	Koe	ssler, J-L
X : parti Y : parti docu	ATEGORY OF CITED DOCUMENTS cularly relevant if taken alone cularly relevant if comb ned with another ment of the same category nological background	T : theory or principle E : earlier patent doc after the filling dat , D : document cited in L : document cited fo	underlying the in ument, but publis	vention hed on, or



Application Number EP 02 00 5113

Category	Citation of document with in of relevant passa	ndication, where appropriate, ages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CI.7)
X	ELECTROPHORESCENT D TRIS(2-PHENYLPYRIDI ELECTRON-TRANSPORTI APPLIED PHYSICS LET INSTITUTE OF PHYSIC vol. 77, no. 6,	NE)IRIDIUM DOPED INTO NG MATERIALS" TERS, AMERICAN S. NEW YORK, US, -08-07), pages 904-906,	1,2,4,8, 9,11-14	·
X	FOR IRIDIUM COMPLEX ORGANIC LIGHT-EMITT JAPANESE JOURNAL OF	NE) AS HOST MATERIAL ES IN HIGH-EFFICIENCY ING DEVICES" APPLIED PHYSICS, JAPANESE JOURNAL OF KYO, JP, RT 2, 1-08-01), pages 235	1,2,4,8, 9,11-14	TECHNICAL FIELDS SEARCHED (Int.Cl.7)
X		, WEINHEIM, DE, 1-08-02), pages 1077	1,2,4,8, 9,11-14	
	The present search report has	·	<u> </u>	
	Place of search Munich	3 February 2004	Koe	essler, J-L
X : part Y : part docu A : tech	ATEGORY OF CITED DOCUMENTS icularly relevant if taken alone icularly relevant if combined with anot iment of the same category inclogical background written disclosure	T : theory or principl E : earlier patent dor after the filing dat her D : document cited i L : document cited i	e underlying the cument, but publice in the application or other reasons	invention shed on, or



Application Number EP 02 00 5113

Category	Citation of document with it of relevant pass	ndication, where appropriate, ages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CI.7)
x	IN ORGANIC LIGHT-EN IRIDIUM-COMPLEX AS CENTER" JAPANESE JOURNAL OF	APPLIED PHYSICS, JAPANESE JOURNAL OF KYO, JP, PART 2, 1999, pages 108595	Y 1,2,4,8, 9,11-14	
X	WO 00/70655 A (UNIV SOUTHERN CALIFORNIA 23 November 2000 (2 * the whole documer	(US)) 000-11-23)	1,2,4,8, 9,11-14	
X	WO 01/008230 A (UNI CALIFORNIA) 1 Febru * the whole documen	V PRINCETON; UNIV SOUT ary 2001 (2001-02-01)	1,2,4,8, 9,11-14	TECHNICAL FIELDS
X	SOCIETY, EASTON, US	Phosphorescent ium Complexes" , AMERICAN CHEMICAL , pages 1704-1711,	1-4,8-14	SEARCHED (Int.Cl.7)
	The present search report has			
	Place of search Munich	Date of completion of the search 3 February 2004	Koe	ssler, J-L
X : part Y : part docu A : tech	ATEGORY OF CITED DOCUMENTS icularly relevant if taken alone icularly relevant if combined with anotiment of the same category inological background written disclosure	T: theory or princip E: earlier patent d after the filing d her D: document cited L: document cited	ole underlying the ir ocument, but publis ate in the application for other reasons	vention



Application Number EP 02 00 5113

Category	Citation of document with in of relevant passa	dication, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CI.7)
P,X	WANG Y ET AL: "HIG ELECTROLUMINESCENT FLUORINATED ORGANOM COMPOUNDS" APPLIED PHYSICS LET INSTITUTE OF PHYSIC	HLY EFFICIENT MATERIALS BASED ON ETALLIC IRIDIUM TERS, AMERICAN S. NEW YORK, US, July 2001 (2001-07-23), 1077255	1,2,4,8, 9,11-14	
X	PHOSPHORESCENT PT(1 ORGANIC ELECTRONICS NL,	ING DIODES UTILIZING I) AND IR(III) DOPANTS" , ELSEVIER, AMSTERDAM, h 2001 (2001–03), pages	1,2,4,8,9,11-14	
P,X	GRUSHIN V V ET AL: electroluminescent organometallic Ir c CHEMICAL COMMUNICAT CHEMISTRY, GB, 2001 XP002196401 ISSN: 1359-7345 * the whole documen	materials based on omplexes" IONS, ROYAL SOCIETY OF , pages 1494-1495,	1,2,4,8,9,11-14	TECHNICAL FIELDS SEARCHED (Int.Cl.7)
P,X	US 2001/019782 A1 (6 September 2001 (2 * the whole documen	KIMURA KEIZO ET AL) 001-09-06) t * 	1-14	
	The present search report has b			
	Place of search Munich	Date of completion of the search 3 February 2004	Koo	essler, J-L
X : parti Y : parti docu A : tech O : non	ATEGORY OF CITED DOCUMENTS icularly relevant if taken alone cularly relevant if combined with anoth ment of the same category nological background -written disclosure mediate document	T : theory or principle E : earlier patent doc after the filing dat b : document cited in L : document cited for	e underlying the in cument, but publis e n the application or other reasons	nvention shed on, or



Application Number EP 02 00 5113

	DOCUMENTS CONSIDE	RED TO BE RELEVANT		
Category	Citation of document with in- of relevant passa		Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CI.7)
P,X		CT IN ORGANIC NCE EMITTING DEVICES ICALLY HINDERED SPACERS MOLECULES" VCH , WEINHEIM, DE, 1-08-16), pages 233	1,2,4,8, 9,11-14	
P,X	PATENT ABSTRACTS OF vol. 2000, no. 26, 1 July 2002 (2002-0) & JP 2001 257076 A 21 September 2001 (2 * abstract *	7-01) (TDK CORP:KIDO JUNJI).	1,2,4,8, 9,11-14	
P,X	EP 1 138 746 A (SUM 4 October 2001 (200 * the whole documen	1-10-04)	1,2,4,8, 9,11-14	TECHNICAL FIELDS SEARCHED (Int.CI.7)
>,х	WO 01/072927 A (IDE 4 October 2001 (200 * the whole documen	1-10-04)	1,2,4,8, 9,11-14	
> , x	PATENT ABSTRACTS OF vol. 2002, no. 03, 3 April 2002 (2002-0 & JP 2001 313179 A CORP), 9 November 20 * abstract *	04-03) (MITSUBISHI CHEMICALS	1,2,4,8,9,11-14	
P,X	US 2001/053462 A1 (I 20 December 2001 (20 * the whole documen	901-12-20)	1,2,4,8, 9,11-14	
	The present search report has b	een drawn up for all claims		
	Place of search	Date of completion of the search	J	Examiner
	Munich	3 February 2004	Koe	ssler, J-L
X : parti Y : parti docu A : tech O : non-	ATEGORY OF CITED DOCUMENTS icularly relevant if taken alone icularly relevant if combined with anotherment of the same category nological background written disclosure mediate document	T: theory or principle E: earlier patent doc after the filing dat er D: document cited in L: document cited for 8: member of the sa document	rument, but publis e n the application or other reasons	shed on, or

EPO FORM 1503 03.82 (P04C01)



Application Number EP 02 00 5113

	DOCUMENTS CONSIDE	RED TO BE RELEVANT		
Category	Citation of document with income of relevant passa		Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.7)
P,X	WO 01/091203 A (AKI KORO (JP); SHOWA DEI 29 November 2001 (20 * the whole documen	NKO KK (JP)) 901-11-29)	1,2,4,8, 9,11-14	
P,X	EP 1 160 889 A (SEM: 5 December 2001 (200 * the whole documen		1,2,4,8, 9,11-14	
P,X	PATENT ABSTRACTS OF vol. 2002, no. 04, 4 August 2002 (2002 & JP 2001 357977 A LTD), 26 December 20 * abstract *	-08-04) (FUJI PHOTO FILM CO	1,2,4,8, 9,11-14	
x	COLOMBO ET AL.: INORG. CHEM., vol. 33, 1994, page: * page 549; table 1	s 545-550, XP002255816	1,2,4,8,9,11-14	
				TECHNICAL FIELDS SEARCHED (Int.Cl.7)
	The present search report has b	een drawn up for all claims		
	Place of search	Date of completion of the search	'	Examiner
	Munich	3 February 2004	Koe	essler, J-L
X : parti Y : parti docu A : tech	ATEGORY OF CITED DOCUMENTS cularly relevant if taken alone cularly relevant if combined with anoth ment of the same category nological background	L : document cited	ocument, but publists ite in the application for other reasons	
	-written disclosure mediate document	& ; member of the s document	same patent tamily	r, corresponding



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CLAIMS INCURRING FEES
The present European patent application comprised at the time of filing more than ten claims.
Only part of the claims have been paid within the prescribed time limit. The present European search report has been drawn up for the first ten claims and for those claims for which claims fees have been paid, namely claim(s):
No claims fees have been paid within the prescribed time limit. The present European search report has been drawn up for the first ten claims.
LACK OF UNITY OF INVENTION
The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:
see sheet B
All further search fees have been paid within the fixed time limit. The present European search report has been drawn up for all claims.
As all searchable claims could be searched without effort justifying an additional fee, the Search Division did not invite payment of any additional fee.
Only part of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the inventions in respect of which search fees have been paid, namely claims:
None of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the invention first mentioned in the claims, namely claims:



LACK OF UNITY OF INVENTION SHEET B

Application Number

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The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:

1. claims: 2,4,9,11; partially 1,5,6-8,12-14

Metal coordination compound of formula (1) wherein ML'n is represented by formula (3), electroluminescent device comprising such a coordination compound and picture display comprising the device.

2. claims: 3,10; partially 1,5-8,12-14

Metal coordination compound of formula (1) wherein ML'n is represented by formula (4), electroluminescent device comprising such a coordination compound and picture display comprising the device.

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 02 00 5113

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

03-02-2004

Patent document cited in search report		Publication date		Patent family member(s)	,	Publication date
EP 1191612	A	27-03-2002	JP EP US	2003081988 1191612 2002063516	A2	19-03-200 27-03-200 30-05-200
EP 1191613	Α	27-03-2002	JP EP US	2003146996 1191613 2002064681	A2	21-05-200 27-03-200 30-05-200
EP 1175128	A	23-01-2002	EP JP US	1175128 2002100476 2002028329	Α	23-01-200 05-04-200 07-03-200
WO 0202714	A	10-01-2002	AU CA CN EP JP WO WO US US	7155001 2411624 1449640 1295514 2004503059 0202714 03063555 2002190250 2003197183 2002121638	A1 T A2 T A2 A1 A1 A1	14-01-200 10-01-200 15-10-200 26-03-200 29-01-200 10-01-200 31-07-200 19-12-200 23-10-200
WO 0215645	A	21-02-2002	AU CN EP WO US	8327401 1454448 1325671 0215645 2002182441	T A1 A1	25-02-20 05-11-20 09-07-20 21-02-20 05-12-20
EP 1211257	A	05-06-2002	JP CN EP US	2003081989 1364847 1211257 2003054198	A A2	19-03-200 21-08-200 05-06-200 20-03-200
WO 0245466	A	06-06-2002	AU EP WO WO US US EP	2256502 2256602 1348711 0245466 0244189 2003059646 2003068526 1349435	A A1 A1 A1 A1 A1	11-06-200 11-06-200 01-10-200 06-06-200 06-06-200 27-03-200 10-04-200 01-10-200
WO 03000661	Α	03-01-2003	JP WO	2003007469 03000661		10-01-200 03-01-200

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 02 00 5113

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

03-02-2004

	Patent document ed in search report		Publication date		Patent family member(s)		Publication date
WO	0141512	A	0 7-06-2001	AU CN EP JP WO US	1807201 1413426 1252803 2003515897 0141512 2003017361 2002034656	T A1 T A1 A1	12-06-2 23-04-2 30-10-2 07-05-2 07-06-2 23-01-2 21-03-2
WO	0070655	Α	23-11-2000	AU BR JP TW WO US US	5004700 0010424 2003526876 500787 0070655 2003017361 2002034656	A T B A2 A1	05-12-2 13-02-2 09-09-2 01-09-2 23-11-2 23-01-2
WO	0108230	A	01-02-2001	US AU CN EP JP WO US	6310360 6113800 1402885 1204994 2003520391 0108230 2003178619 2002008233	A T A1 T A1 A1	30-10-20 13-02-20 12-03-20 15-05-20 02-07-20 01-02-20 25-09-20 24-01-20
US	2001019782	A1	06-09-2001	JP JP	2001345183 2001247859		14-12-2 14-09-2
JP	2001257076	A	21-09-2001	NONE			
EΡ	1138746	Α	04-10-2001	EP JP US	1138746 2001342459 2002027623	A	04-10-2 14-12-2 07-03-2
WO	0172927	A	04-10-2001	CN EP WO TW US	1365381 1205527 0172927 532048 2002045061	A1 A1 B	21-08-20 15-05-20 04-10-20 11-05-20 18-04-20
JP	2001313179	Α	09-11-2001	NONE			
	2001053462	A1	20-12-2001	JР	2001319780		16-11-2
WO	0191203	Α	29-11-2001	AU CA	5678101 2380067	A A1	03-12-2 29-11-2

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 02 00 5113

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

03-02-2004

Patent document cited in search report		Publication date		Patent family member(s)		Publication date
WO 0191203	A		EP WO JP TW US	1214746 0191203 2002050483 518768 2002146589	A2 A B	19-06-200 29-11-200 15-02-200 21-01-200 10-10-200
EP 1160889	A	05-12-2001	CN EP JP TW US	1325143 1160889 2002050484 536836 2001050373	A2 A B	05-12-200 05-12-200 15-02-200 11-06-200 13-12-200
JP 2001357977	Α	26-12-2001	NONE			
						1

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